

# STIC Search Report Biotech-Chem Library

### STIC Database Tracking Number: 133401

TO: Shailendra Kumar Location: 5c03 / 5c18

Wednesday, September 29, 2004

Art Unit: 1521 Phone: 272-0640

Serial Number: 10 / 680979

From: Jan Delaval

**Location: Biotech-Chem Library** 

**Rem 1A51** 

Phone: 272-2504

jan.delaval@uspto.gov

# Search Notes



Aro Deans

PTO-1590 (8-01)

## 133401

Access	DB#	

### SEARCH REQUEST FORM

### Scientific and Technical Information Center

Requester's Fufl Name:  Art Unit:  Mail Box and Bldg/Room Location	n: <u>NEW 5003</u> Res 5018	sults Format Preferred (circle): PA	e: (1) 73 ) 04 EC 979 PER DISK E-MAIL
If more than one search is subn ****************************  Please provide a detailed statement of the Include the elected species or structures, I utility of the invention. Define any terms known. Please attach a copy of the cover	search topic, and describe keywords, synonyms, acre that may have a special n sheet, pertinent claims. an	*************************  e as specifically as possible the subject monyms, and registry numbers, and combineaning. Give examples or relevant citated abstract.	atter to be searched, the with the concept or tions, authors, etc, if
Title of Invention: $\sqrt{r \approx c \leq s} = \epsilon$ Inventors (please provide full names):	Yong-Mic	on Chai et al	els his the officer is
Earliest Priority Filing Date: *For Sequence Searches Only* Please inclu	, ,		numbers) along with the
appropriate serial number.	ае ан регипен ипјогтанов	(parent, chua, atvistonat, or issuea patent i	umbers) along with the
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Sec claims 1, 8,	9, 10, 111	7.4	
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Searcher Location:	Structure (#)	Questel/Orbit	
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Date Completed: 5/29	Litigation		
Searcher Prep & Review Time:	Fulltext	C. 7 CO.	
Clerical Prep Time: 30	Patent Family	WWW/Internet Other (specify)	*
Online Time:	Other	Other (specify)	· · · · ·

=> fil casreact FILE 'CASREACT' ENTERED AT 10:03:38 ON 29 SEP 2004 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE CONTENT: 1840 - 26 Sep 2004 VOL 141 ISS 13

Some CASREACT records are derived from the ZIC/VINITI database (1974-1991) provided by InfoChem, INPI data prior to 1986, and Biotransformations database compiled under the direction of Professor Dr. Klaus Kieslich.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d sta que 124 L18 STR

 $N \sim C \sim G1 \sim C$   $1 \quad 2 \quad 3 \quad 4$ 

REP G1=(0-5) CH2 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 4

STEREO ATTRIBUTES: NONE L21 STR

RRT PRO 13

N \( \infty \) C - G1 - C - OH

1 2 3 4 5

N \( \infty \) C - G1 - C - O - C - N

6 7 8 9 10 11 12

REP G1=(0-5) CH2 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 13

STEREO ATTRIBUTES: NONE

L23 233 SEA FILE=CASREACT SSS FUL L21 ( 1681 REACTIONS)

L24 233 SEA FILE=CASREACT SUB=L23 SSS FUL L18 ( 1681 REACTIONS)

1681 HIT RXNS

233 DOCS

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                SET COST OFF
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            442 S E3, E17
L1
                E CHOI YONG/AU
            129 S E3, E60, E63
L2
                E CHOI YOUNGMOON/AU
              1 S E3
L3
                E KIM M/AU
            264 S E3, E30, E31
L4
                E KIM MIN/AU
             79 S E3, E98
L5
             5 S E148
L6
                E SK/PA,CS
             85 S E45-E68
L7
          27657 S E3,E4
L8
              6 S O CARBAMOYL AND L1-L8
Ь9
              7 S O CARBAM? AND L1-L8
L10
              7 S L10, L9
L11
              8 S L1-L3 AND L4-L6
L12
              6 S L1-L6 AND L7, L8
L13
             14 S L12, L13
L14
             13 S L14 NOT L11
L15
              6 S L15 AND ?CARBAM?
L16
L17
             13 S L11, L16
     FILE 'CASREACT' ENTERED AT 07:53:52 ON 29 SEP 2004
                STR
L18
                STR L18
L19
L20
              0 S L19
                STR L19
L21
L22
              7 S L21
            233 S L21 FUL
L23
                SAV TEMP L23 KUMAR680/A
            233 S L18 FUL SUB=L23
L24
                SAV TEMP L24 KUMAR680A/A
     FILE 'REGISTRY' ENTERED AT 08:31:10 ON 29 SEP 2004
              5 S (SODIUM CYANATE OR POTASSIUM CYANATE OR AMMONIUM CYANATE OR M
L25
              1 S 420-05-3
L26
            235 S 420-05-3/CRN
L27
             12 S (HYDROCHLORIC ACID OR SULFURIC ACID OR PHOSPHORIC ACID OR ACE
L28
             12 S (DICHLOROMETHANE OR ACETONITRILE OR CHLOROFORM OR 1,2-DICHLOR
L29
     FILE 'HCAPLUS' ENTERED AT 08:34:57 ON 29 SEP 2004
     FILE 'CASREACT' ENTERED AT 08:35:03 ON 29 SEP 2004
              3 S L25, L26 AND L24
L30
L31
              4 S L27 AND L24
             89 S L28 AND L24
L32
            135 S L29 AND L24
L33
              3 S L30, L31 AND L32, L33
L34
```

4 S L30, L31, L34

4 S L35 AND L23

L35 L36

```
FILE 'REGISTRY' ENTERED AT 08:38:39 ON 29 SEP 2004
L37
                STR
             50 S L37
L38
                STR L37
L39
                STR L39
L40
             50 S L40
L41
          23281 S L40 FUL
L42
     FILE 'HCAPLUS' ENTERED AT 08:41:48 ON 29 SEP 2004
           3187 S L42(L) PREP+NT/RL OR L42/P
L43
L44
          16517 S L42
L45
           2794 S L25, L26
           2029 S (NA OR SODIUM OR K OR POTASSIUM OR NH3 OR AMMONIUM OR MG OR M
L46
             13 S L43 AND L45, L46
L47
L48
             11 S (NAOCN OR KOCN OR NH3OCN OR MGOCN OR CAOCN) AND L43
L49
             0 S (NA OR K OR NH3 OR MG OR CA) () OCN AND L43
L50
             13 S L44 AND L45, L46
             14 S (NAOCN OR KOCN OR NH3OCN OR MGOCN OR CAOCN) AND L44
L51
             1 S (NA OR K OR NH3 OR MG OR CA) () OCN AND L44
L52
             24 S L47, L48, L50-L52
L53
             12 S L43 AND L27
L54
             12 S L44 AND L27
L55
             25 S L53-L55
L56
            34 S L28 AND L43
L57
L58
            137 S L28 AND L44
L59
             33 S L29 AND L43
L60
             79 S L29 AND L44
L61
              3 S L56 AND L57-L60
L62
            226 S L47-L61
L63
           3103 S L43 NOT L62
     FILE 'REGISTRY' ENTERED AT 08:48:04 ON 29 SEP 2004
     FILE 'HCAPLUS' ENTERED AT 08:48:04 ON 29 SEP 2004
                SET SMARTSELECT ON
            SEL L62 1- RN : 20116 TERMS
L64
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 08:48:23 ON 29 SEP 2004
        20116 S L64
L65
     FILE 'HCAPLUS' ENTERED AT 08:49:49 ON 29 SEP 2004
                SET SMARTSELECT ON
L66
            SEL L63 1- RN : 50645 TERMS
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 08:50:19 ON 29 SEP 2004
L67
          50625 S L66
     FILE 'HCAPLUS' ENTERED AT 08:53:12 ON 29 SEP 2004
L68
           3103 S L63 OR L63
L69
            500 S L68 RAN=(2001:713296,)
            500 S L68 RAN=(1998:265055,2001:704856)
L70
            500 S L68 RAN=(1993:537527,1998:263478)
L71
L72
            500 S L68 RAN=(1986:568756,1993:534388)
            500 S L68 RAN=(1976:592613,1986:552768)
L73
L74
            603 S L68 RAN=(,1976:592573)
     FILE 'REGISTRY' ENTERED AT 08:55:29 ON 29 SEP 2004
```

FILE 'HCAPLUS' ENTERED AT 08:55:30 ON 29 SEP 2004

SET SMARTSELECT ON

L75 SEL L74 1- RN : 12729 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 08:55:52 ON 29 SEP 2004 L76 12729 S L75

SET SMARTSELECT ON SET SMARTSELECT OFF

FILE 'HCAPLUS' ENTERED AT 08:57:24 ON 29 SEP 2004 SET SMARTSELECT ON

L77 SEL L73 1- RN : 24463 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 08:57:43 ON 29 SEP 2004 L78 24462 S L77

FILE 'HCAPLUS' ENTERED AT 08:58:56 ON 29 SEP 2004 SET SMARTSELECT ON

L79 SEL L72 1- RN : 35353 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 08:59:30 ON 29 SEP 2004 L80 35353 S L79

FILE 'HCAPLUS' ENTERED AT 09:01:14 ON 29 SEP 2004 SET SMARTSELECT ON

L81 SEL L71 1- RN : 42897 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 09:02:05 ON 29 SEP 2004 L82 42897 S L81

FILE 'HCAPLUS' ENTERED AT 09:04:38 ON 29 SEP 2004 SET SMARTSELECT ON

L83 SEL L70 1- RN : 50471 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 09:05:24 ON 29 SEP 2004 L84 50471 S L83

FILE 'HCAPLUS' ENTERED AT 09:08:39 ON 29 SEP 2004

L85 250 S L68 RAN=(2003:173580,)

L86 250 S L68 RAN=(2001:713296,2003:168865)

L87 250 S L68 RAN=(2000:43347,2001:704856)

L88 250 S L70 NOT L87

FILE 'REGISTRY' ENTERED AT 09:13:06 ON 29 SEP 2004

FILE 'HCAPLUS' ENTERED AT 09:13:06 ON 29 SEP 2004

SET SMARTSELECT ON

L89 SEL L88 1- RN : 26803 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 09:13:24 ON 29 SEP 2004 L90 26803 S L89

FILE 'HCAPLUS' ENTERED AT 09:14:59 ON 29 SEP 2004 SET SMARTSELECT ON

L91 SEL L87 1- RN : 36295 TERMS SET SMARTSELECT OFF

FILE 'REGISTRY' ENTERED AT 09:15:27 ON 29 SEP 2004 L92 36295 S L91

```
FILE 'HCAPLUS' ENTERED AT 09:17:49 ON 29 SEP 2004
                SET SMARTSELECT ON
            SEL L86 1- RN : 41291 TERMS
L93
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 09:18:08 ON 29 SEP 2004
L94
          41291 S L93
     FILE 'HCAPLUS' ENTERED AT 09:20:58 ON 29 SEP 2004
                SET SMARTSELECT ON
            SEL L85 1- RN : 50645 TERMS
L95
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 09:21:30 ON 29 SEP 2004
          50645 S L95
L96
     FILE 'HCAPLUS' ENTERED AT 09:24:09 ON 29 SEP 2004
L97
            125 S L85 RAN=(2003:841484,)
            125 S L85 NOT L97
L98
     FILE 'REGISTRY' ENTERED AT 09:24:55 ON 29 SEP 2004
     FILE 'HCAPLUS' ENTERED AT 09:24:55 ON 29 SEP 2004
                SET SMARTSELECT ON
            SEL L98 1- RN : 23914 TERMS
L99
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 09:25:09 ON 29 SEP 2004
          23914 S L99
L100
     FILE 'HCAPLUS' ENTERED AT 09:26:50 ON 29 SEP 2004
                SET SMARTSELECT ON
                              32103 TERMS
L101
            SEL L97 1- RN :
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 09:27:05 ON 29 SEP 2004
L102
         32103 S L101
         239012 S L65,L67,L76,L78,L80,L82,L84,L90,L92,L94,L96,L100,L102
L103
L104
             50 S L39 SAM SUB=L103
          72933 S L39 FUL SUB=L103
L105
                STR L39
L106
                STR L106
L107
          72933 S L105 OR L105
L108
          36000 S L108 RAN=(192723-34-5,)
L109 🕜
          36933 S L108 NOT L109
L110
     FILE 'HCAPLUS' ENTERED AT 09:40:39 ON 29 SEP 2004
        1076566 S L109 OR L110
L111
           177 S L62 AND L111
L112
           2578 S L63 AND L111
L113
             11 S L112, L113 AND L45, L46
L114
             19 S L112, L113 AND L53
L115
             19 S L114, L115
L116
             2 S L116 AND L28, L29
L117
             11 S L111(L) RACT+NT/RL AND L116
L118
             70 S L111(L) RACT+NT/RL AND L112
L119
L120
             11 S L117, L118
             8 S L1-L8 AND L43
L121
L122
             26 S L1-L8 AND L44
L123 -
             12 S L121, L122 AND L111
             7 S L123 AND L112,L113
L124
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L125

18 S L120, L124

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5 S L123 NOT L125
L126
            18 S L125 AND (PD<=20031008 OR PRD<=20031008 OR AD<=20031008)
L127
     FILE 'REGISTRY' ENTERED AT 09:45:26 ON 29 SEP 2004
          2505 S L42 AND 46.150.18/RID AND 1/NR
L128
            13 S L128 AND C10H14N2O2
L129
               SEL RN 1 2 9 10 11 12 13
              6 S L129 NOT E1-E7
L130
           3672 S L105 AND 46.150.18/RID AND 1/NR
L131
             22 S L131 AND C9H13NO
L132
               SEL RN 14 17 16 6
              4 S E8-E11
L133
     FILE 'HCAPLUS' ENTERED AT 09:50:02 ON 29 SEP 2004
             1 S L130
L134
              1 S L133 AND L134
L135
     FILE 'REGISTRY' ENTERED AT 09:50:17 ON 29 SEP 2004
             55 S L42 AND NC5-C6/ES AND 2/NR
L136
             6 S L136 AND C11H14N2O2
L137
            277 S L105 AND NC5-C6/ES AND 2/NR
L138
              5 S L138 AND C10H13NO
L139
               SEL RN 3 5
L140
              3 S L139 NOT E12-E13
     FILE 'HCAPLUS' ENTERED AT 09:52:27 ON 29 SEP 2004
              1 S L137
L141
L142
              1 S L141 AND L140
    FILE 'REGISTRY' ENTERED AT 09:52:47 ON 29 SEP 2004
            996 S L42 AND 46.150.18/RID AND NC5/ES AND 3/NR
L143
              6 S L143 AND C21H23FN2O3
L144
           3756 S L105 AND 46.150.18/RID AND NC5/ES AND 3/NR
L145
             0 S L145 AND C20H22FNO2
             40 S C20H22FNO2 AND 46.150.18/RID AND NC5/ES AND 3/NR
L147
              6 S L147 AND METHANONE AND 4 FLUOROPHENYL AND 2 HYDROXY 2 PHENYLE
L148
    FILE 'HCAPLUS' ENTERED AT 09:57:39 ON 29 SEP 2004
              1 S L144
L149
              0 S L149 AND L148
L150
L151
             3 S L148
             22 S L127, L135, L142, L149, L151
L152
             11 S L152 AND ?CYANAT?
L153
             11 S L152 AND L45, L46
L154
             12 S L153, L154
L155
             10 S L152 NOT L155
L156
              5 S L156 AND L135, L142, L149, L151
L157
              5 S L156 NOT L157
L158
     FILE 'CASREACT' ENTERED AT 10:00:55 ON 29 SEP 2004
     FILE 'CASREACT' ENTERED AT 10:03:38 ON 29 SEP 2004
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L36 ANSWER 1 OF 4 CASREACT COPYRIGHT 2004 ACS on STN

AN 137:124782 CASREACT

TI Method for carbamoylating alcohols with an alkali metal cyanate in the presence of methanesulfonic, sulfuric or acetic acids

IN Ellis, James E.

PA USA

SO U.S. Pat. Appl. Publ., 8 pp.

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CODEN: USXXCO
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DTPatent English LA

PAT	ENT 1	10.				DATE								DATE			
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								US 2002-56268				20020125					
US	66139	908		B2	2	2003	0902										
WO	20020	06089	93	A:	1	2002	8080		W	200	02-II	382		20020	)111		
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		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KΡ,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,
		UG,	US,	UZ,	VN,	ΥU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	ΤZ,	ŪĠ,	ZM,	ZW,	ΑT,	BE,	CH,
		CY,	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,
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US	2001	-265	502P	200	0101	31											
WO	2002	-IB82	2	20	0201	11											
	US US WO	US 2002: US 6613: WO 2002: W: RW: EP 1377: R: BR 2002: JP 2004: US 2001:	PATENT NO.  US 20021033 US 6613908 W: AE, CO, GM, LS, PL, UG, RW: GH, CY, BF, EP 1377568 R: AT, IE, BR 20020068 US 2001-265	PATENT NO.	PATENT NO. KIN  US 2002103378 A:  US 6613908 B:  WO 2002060893 A:  W: AE, AG, AL,  CO, CR, CU,  GM, HR, HU,  LS, LT, LU,  PL, PT, RO,  UG, US, UZ,  RW: GH, GM, KE,  CY, DE, DK,  BF, BJ, CF,  EP 1377568 A:  R: AT, BE, CH,  IE, SI, LT,  BR 2002006806 A  JP 2004518687 T:  US 2001-265502P 206	PATENT NO. KIND	PATENT NO. KIND DATE  US 2002103378 A1 2002  US 6613908 B2 2003  W: AE, AG, AL, AM, AT,  CO, CR, CU, CZ, DE,  GM, HR, HU, ID, IL,  LS, LT, LU, LV, MA,  PL, PT, RO, RU, SD,  UG, US, UZ, VN, YU,  RW: GH, GM, KE, LS, MW,  CY, DE, DK, ES, FI,  BF, BJ, CF, CG, CI,  EP 1377568 A1 2004  R: AT, BE, CH, DE, DK,  IE, SI, LT, LV, FI,  BR 2002006806 A 2004	PATENT NO. KIND DATE  US 2002103378 A1 20020801  US 6613908 B2 20030902  WO 2002060893 A1 20020808  W: AE, AG, AL, AM, AT, AU, CO, CR, CU, CZ, DE, DK, GM, HR, HU, ID, IL, IN, LS, LT, LU, LV, MA, MD, PL, PT, RO, RU, SD, SE, UG, US, UZ, VN, YU, ZA, RW: GH, GM, KE, LS, MW, MZ, CY, DE, DK, ES, FI, FR, BF, BJ, CF, CG, CI, CM, EP 1377568 A1 20040107  R: AT, BE, CH, DE, DK, ES, IE, SI, LT, LV, FI, RO, BR 2002006806 A 20040203 JP 2004518687 T2 20040624	PATENT NO. KIND DATE  US 2002103378 A1 20020801  US 6613908 B2 20030902  WO 2002060893 A1 20020808  W: AE, AG, AL, AM, AT, AU, AZ, CO, CR, CU, CZ, DE, DK, DM, GM, HR, HU, ID, IL, IN, IS, LS, LT, LU, LV, MA, MD, MG, PL, PT, RO, RU, SD, SE, SG, UG, US, UZ, VN, YU, ZA, ZW, RW: GH, GM, KE, LS, MW, MZ, SD, CY, DE, DK, ES, FI, FR, GB, BF, BJ, CF, CG, CI, CM, GA, EP 1377568 A1 20040107  R: AT, BE, CH, DE, DK, ES, FR, IE, SI, LT, LV, FI, RO, MK, BR 2002006806 A 20040203 JP 2004518687 T2 20040624 US 2001-265502P 20010131	PATENT NO. KIND DATE AND COLORS A	PATENT NO. KIND DATE APPLICATION APPLICATI	PATENT NO. KIND DATE APPLICATION OF THE PRICE OF THE PRIC	PATENT NO. KIND DATE APPLICATION NO.  US 2002103378 A1 20020801 US 2002-56268  US 6613908 B2 20030902  WO 2002060893 A1 20020808 WO 2002-IB82  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, EP 1377568 A1 20040107 EP 2002-73761: R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR  BR 2002006806 A 20040203 BR 2002-6806 JP 2004518687 T2 20040624 JP 2002-56104: US 2001-265502P 20010131	PATENT NO. KIND DATE APPLICATION NO.  US 2002103378 A1 20020801 US 2002-56268  US 6613908 B2 20030902  WO 2002060893 A1 20020808 WO 2002-IB82  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, EP 1377568 A1 20040107 EP 2002-737611  R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, TE, SI, LT, LV, FI, RO, MK, CY, AL, TR  BR 2002006806 A 20040203 BR 2002-6806  JP 2004518687 T2 20040624 JP 2002-561042  US 2001-265502P 20010131	PATENT NO. KIND DATE APPLICATION NO. DATE  US 2002103378 A1 20020801 US 2002-56268 20020  WO 2002060893 A1 20020808 WO 2002-IB82 20020  W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, EP 1377568 A1 20040107 EP 2002-737611 20020  R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR  BR 2002006806 A 20040203 BR 2002-6806 20020  US 2001-265502P 20010131	PATENT NO.	PATENT NO.

The present invention includes a method for carbamoylating an alc. with AB sodium cyanate in the presence of methanesulfonic acid. The reaction can be conducted under anhydrous conditions. This method is suitable for carbamoylating a mol. including both an alc. moiety and a basic moiety and/or a mol. including both an alc. moiety and a sulfenyl moiety, such as the sulfenyl alc. precursor of the antiviral agent Capravirine.

### RX(1) OF 1

C YIELD 95%

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RX(1) RCT A 917-61-3, B 178981-89-0
```

STAGE (1)

CAT **75-75-2** MeSO3H SOL **75-05-8** MeCN

STAGE (2)

PRAI GB 2000-24811

WO 2001-EP11603 20011008

SOL 7732-18-5 Water

20001010

PRO C **178979-85-6** 

NTE optimization study

```
ANSWER 2 OF 4 CASREACT COPYRIGHT 2004 ACS on STN
     136:325706 CASREACT
     Preparation of pleuromutilin derivatives as antibacterial agents
TI
     Elder, John Stephen; Forrest, Andrew Keith; Jarvest, Richard Lewis;
IN
     Sheppard, Robert John
PA
     Smithkline Beecham P.L.C., UK
     PCT Int. Appl., 54 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
                                 DATE
                         KIND
                                                   APPLICATION NO.
     PATENT NO.
                                 _____
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                                 20020418
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PΙ
     WO 2002030929
                         A1
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                                                                            TZ, UA,
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     AU 2002018215
                                 20020422
                                                   AU 2002-18215
                                                                       20011008
                           Α5
     EP 1351959
                           A1
                                 20031015
                                                   EP 2001-986687
                                                                       20011008
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                                                       20011008
     JP 2004511482
                           T2
                                 20040415
                                                   JP 2002-534315
                                                   US 2003-399023
                                                                       20030725
     US 2004024059
                                 20040205
                           A1
```

$$R^2$$
 Me OH  $R^2$  Me  $R$ 

Pleuromutilin derivs., such as I or II [R1 = (substituted) heterocycle, AB alkyl, cycloalkyl, heteroaryl; R2= vinyl, Et; R3 = H, OH, F; R4 = H, F; R5, R6 = H, OH; R5R6 = oxo], were prepared for the use in antibacterial therapy. Thus, reaction between 2-(methylsulfonyl)ethyl chloroformate and (3R) -3-deoxo-11-deoxy-3-methoxy-11-oxo-4-epimutilin provided (3R) -3-deoxo-11-deoxy-3-methoxy-11-oxo-4-epimutilin 14-[N-(2-1)]methylsulfonylethoxycarbonyl)]carbamate, which on selective oxidation of 3-methoxyl group and simultaneous reduction of 11-oxo group, afforded pleuromutilin derivative I [R1 = CH2CH2SO2Me; R2 = CH:CH2; R3, R4 = H; R5R6 = O (III)]. The prepared pleuromutilin derivs. were tested for antibacterial activity against Staphylococcus aureus Oxford, Streptococcus pneumoniae 1629, Moraxella catarrhalis 502 and Haemophilus influenzae Q1, e.g. III MIC =  $\leq 4 \mu q/mL$  (S. aureus).

### RX(2) OF 124 D

YIELD 21%

RCT D 412278-62-7, E 3315-16-0 RX(2)

STAGE(1)

RGT H 110-86-1 Pyridine SOL 75-09-2 CH2Cl2

STAGE(2)

RCT F 67-56-1

STAGE(3)

RGT I 7647-01-0 HCl SOL 123-91-1 Dioxane

PRO G 412275-39-9

RETABLE

Referenced Author (RAU)	(RPY) (	•	Referenced Work (RWK)	Referenced   File
=======================================	+====+=	====+		
Brooks, G	2001		WO 0174788 A	CAPLUS
Dominic, S	11999	i	WO 9921855 A	CAPLUS
DOMITITE, 5	1 =	!		CA DI IIG
Hunt, E	1997	ł	WO 9725309 A	CAPLUS
Naylor, A	1998	İ	WO 9805659 A	CAPLUS
Naylor, A	2001	İ	WO 0114310 A	CAPLUS

ANSWER 3 OF 4 CASREACT COPYRIGHT 2004 ACS on STN L36

106:130551 CASREACT AN

Synthesis and reactivities of triisocyanatoantimony ΤI

Kijima, Ichiro; Wakeshima, Ikuko; Sasaki, Toru UΑ

Fac. Eng., Sci. Univ. Tokyo, Tokyo, 162, Japan CS

Nippon Kagaku Kaishi (1986), (12), 1754-57

CODEN: NKAKB8; ISSN: 0369-4577

DT Journal

so

Japanese LΑ

Sb(NCO)3 was prepared by the reaction of SbCl3 with NaOCN in the presence of AB several additives in benzene and THF. The reaction was accelerated remarkably by using THF as an additive in benzene to give Sb(NCO)3 in high yield. Sb(NCO)3 reacted with amines such as NHEt2, BuNH2, PhH2, and NH3 to afford only the corresponding triureidoantimony compds., but reacted with alcs. such as iso-PrOH, BuOH, sec- and tert-BuOH or PhOH to yield the corresponding carbamate and trialkoxo- or triphenoxyantimony compds. Sb(NCO)3 reacted also with 2-diethylaminoethanol (HL) to give SbL3 and 2-diethylaminoethyl carbamate, together with isocyanuric acid. Sb(NCO)3 reacted with alcs. and PhOH to yield the corresponding substituted

YIELD 61%

products, but the reaction with amines provided only the corresponding addition products.

P(X) = P(X) = P(X) = P(X) P(X) = P(X) = P(X)

RX(2) RCT D **917-61-3**, B 67-63-0 RGT E 10025-91-9 SbCl3 PRO C 1746-77-6

В

D

ANSWER 4 OF 4 CASREACT COPYRIGHT 2004 ACS on STN L36 106:17586 CASREACT ANAsymmetric syntheses and potential asymmetric synthesis of  $\alpha$ -amino TIalcohols: hydroxyamination of olefins by the sharpless method Ben Hassine, B.; Gorsane, M.; Pecher, J.; Martin, R. H. ΑU Lab. Synth. Org. Photochim., Fac. Sci. Tech., Monastir, 5000, Tunisia CS Bulletin des Societes Chimiques Belges (1985), 94(11-12), 759-69 SO CODEN: BSCBAG; ISSN: 0037-9646 DTJournal LA French GΙ

Optically active  $\alpha$ -amino alcs. were synthesized by the Sharpless method using (-)-10,11-dihydroquinine (I) and (R)-(-)-pantolactone as chiral inducers (R1OH). (dl)-2-Hydroxyheptahelicene (II) and 5 secondary (dl) alcs. were also used to prepare the intermediate diastereomeric (dl) $\alpha$ -hydroxy carbamates III. The highest inductions (e.e.  $\geq 98\%$ ) were obtained with (E)-stilbene and I or II.

RX(1) OF 57 A + B ===> C...

RX(1) RCT A 108-86-1, B 98-03-3 RGT D 7439-95-4 Mg PRO C 26059-21-2 SOL 109-99-9 THF

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 27 SEP 2004 HIGHEST RN 752974-11-1 DICTIONARY FILE UPDATES: 27 SEP 2004 HIGHEST RN 752974-11-1

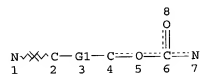
TSCA INFORMATION NOW CURRENT THROUGH MAY 21, 2004

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> d sta que 142 L40 S'



REP G1=(0-5) CH2 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 8

STEREO ATTRIBUTES: NONE L42 23281 SEA FILE=REGISTRY SSS FUL L40

100.0% PROCESSED 224298 ITERATIONS SEARCH TIME: 00.00.02

23281 ANSWERS

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Starting

REP G1=(0-5) CH2 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 5

STEREO ATTRIBUTES: NONE

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FILE COVERS 1907 - 29 Sep 2004 VOL 141 ISS 14 FILE LAST UPDATED: 28 Sep 2004 (20040928/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L155 ANSWER 1 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
    2002:575780 HCAPLUS
    137:124782
    Entered STN: 02 Aug 2002
ED
    Method for carbamoylating alcohols with an alkali metal cyanate
    in the presence of methanesulfonic, sulfuric or acetic acids
    Ellis, James E.
IN
PA
    USA
    U.S. Pat. Appl. Publ., 8 pp.
SO
    CODEN: USXXCO
DT
    Patent
    English
LA
    ICM C07D041-02
     ICS C07C269-00; C07H013-00
NCL
    546272100
    21-2 (General Organic Chemistry)
     Section cross-reference(s): 63
FAN.CNT 1
    PATENT NO.
                        KIND
                               DATE
                                          APPLICATION NO.
                        ____
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-----\_\_\_\_\_ \_\_\_\_\_\_ A1 20020801 US 2002-56268 20020125 <--US 2002103378 PΤ B2 A1 US 6613908 20030902 WO 2002-IB82 20020111 <--20020808 WO 2002060893 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,

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                                           EP 2002-737611
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    BR 2002006806
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                                                                  20020111 <--
                                20040624
     JP 2004518687
PRAI US 2001-265502P
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                                20010131
                                         <--
                         W
                                20020111
    WO 2002-IB82
                                         <--
CLASS
                CLASS PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
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US 2002103378
                ICM
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                        C07C269-00; C07H013-00
                ICS
                        546272100
                NCL
                        C07C269/00; C07D401/06
US 2002103378
                ECLA
                FTERM 4C063/AA01; 4C063/BB03; 4C063/CC25; 4C063/DD12;
 JP 2004518687
                        4C063/EE05; 4H039/CA99; 4H039/CF40
OS
    CASREACT 137:124782
    The present invention includes a method for carbamoylating an alc. with
AB
     sodium cyanate in the presence of methanesulfonic acid.
     The reaction can be conducted under anhydrous conditions. This method is
     suitable for carbamoylating a mol. including both an alc. moiety and a
     basic moiety and/or a mol. including both an alc. moiety and a sulfenyl
     moiety, such as the sulfenyl alc. precursor of the antiviral agent
     Capravirine.
     carbamoylation sulfenyl alc sodium cyanate
ST
     methanesulfonic acid antiviral agent
IT
     Carbamoylation catalysts
        (method for carbamoylating alcs. including sulfenyl alcs. with an
        alkali metal cyanate in the presence of methanesulfonic,
        sulfuric or acetic acids)
     Alcohols, reactions
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (method for carbamoylating alcs. including sulfenyl alcs. with an
        alkali metal cyanate in the presence of methanesulfonic,
        sulfuric or acetic acids)
IT
     Antiviral agents
        (method for carbamoylating alcs. including sulfenyl alcs. with an
        alkali metal cyanate in the presence of methanesulfonic,
        sulfuric or acetic acids suitable for preparation of)
     Heterocyclic compounds
IT
     RL: CAT (Catalyst use); USES (Uses)
        (nitrogen; method for carbamoylating alcs. including sulfenyl alcs.
        with an alkali metal cyanate in the presence of
        methanesulfonic, sulfuric or acetic acids optionally in the presence
        of)
     64-19-7, Acetic acid, uses 75-75-2, Methanesulfonic acid
IT
     7664-93-9, Sulfuric acid, uses
     RL: CAT (Catalyst use); USES (Uses)
        (method for carbamoylating alcs. including sulfenyl alcs. with an
        alkali metal cyanate in the presence of methanesulfonic,
        sulfuric or acetic acids)
     590-28-3, Potassium cyanate 917-61-3
TT
     , Sodium cyanate 21846-90-2, Cesium
     cyanate
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (method for carbamoylating alcs. including sulfenyl alcs. with an
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alkali metal cyanate in the presence of methanesulfonic,
        sulfuric or acetic acids)
     178981-89-0
TT
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (method for carbamoylating alcs. including sulfenyl alcs. with an
        alkali metal cyanate in the presence of methanesulfonic,
        sulfuric or acetic acids suitable for carbamoylation of)
ΙT
     178979-85-6P, Capravirine
     RL: IMF (Industrial manufacture); PUR (Purification or
     recovery); PREP (Preparation)
         (method for carbamoylating alcs. including sulfenyl alcs. with an
        alkali metal cyanate in the presence of methanesulfonic,
         sulfuric or acetic acids suitable for preparation of)
     75-05-8, Acetonitrile, uses 109-99-9, THF, uses
IT
     141-78-6, Ethyl acetate, uses
     RL: NUU (Other use, unclassified); USES (Uses)
         (solvent; method for carbamoylating alcs. including sulfenyl alcs. with
         an alkali metal cyanate in the presence of methanesulfonic,
         sulfuric or acetic acids in)
     64-19-7, Acetic acid, uses
IT
     RL: IMF (Industrial manufacture); PUR (Purification or
     recovery); PREP (Preparation)
         (method for carbamoylating alcs. including sulfenyl alcs. with an
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     64-19-7 HCAPLUS
RN
     Acetic acid (7CI, 8CI, 9CI) (CA INDEX NAME)
CN
HO-C-CH3
L155 ANSWER 2 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
     2002:353634 HCAPLUS
AN
     136:365765
DN
     Entered STN: 12 May 2002
ED
     Inhibitors of transglutaminases
TI
     Fuchsbauer, Hans-Lothar; Pasternack, Ralf; Zotzel, Jens
TN
     N-Zyme Biotec G.m.b.H., Germany
PA
     PCT Int. Appl., 44 pp.
SO
      CODEN: PIXXD2
DT
      Patent
     German
LΑ
      ICM C12P013-00
IC
      7-8 (Enzymes)
      Section cross-reference(s): 34
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      PATENT NO.
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                                                 WO 2001-EP12727
                                                                            20011102 <--
     WO 2002036798
                            A2
                                    20020510
PΙ
     WO 2002036798
                            A3
                                    20030424
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os
     The invention relates to a chemical compound of formula R1(CH2)mYn(CH2)oC(Z)R2
AB
     (I), wherein R1 means formula R4bqNHCH(CH3)C(O)apR3, (II), R6X(CH3)R5, or
     (III); R2 means H, alkyl, which can optionally be substituted with halogen
     or N2, or NH2; m and o mean 0-3 and n means 0 or 1; ap, bq and cr mean
     amino acid chains and p, q, and r mean the number of amino acids, a and/or b
     and/or c also being able to contain ≥1 side chain, represented by
     (CH2) mYn(CH2) oC(Z)R2, Y, Z, R2, m, n, and o having the same meaning as in
     formula I, and p, q and r being the same or different and meaning a whole
     number from 0 to 1000; R3 and R4, independently of each other, mean H, alkyl,
     aryl, a heterocycle, an amino protective group or a carboxy protective
     group; R5 and R6, independently of each other, mean alkyl which can
     contain ≥1 hetero atom selected from N, O and S; aryl or a
     heterocycle; X means a methine group, a N or a P atom; Y means an O atom,
     a S atom or an NH-group; and Z means an O atom, a S atom or an NR7-group,
     R7 meaning H, alkyl, aryl, a heterocycle, O-alkyl, O-aryl, O-heterocycle,
     NR2 or NHCONR2, R meaning H, alkyl, aryl or a heterocycle; and to the use
     of said compound as an inhibitor of transglutaminases.
     transqlutaminase inhibitor
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        (N, N-Di-Me; inhibitors of transglutaminases)
     Caseins, biological studies
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     590-28-3, Potassium cyanate
     Pentafluorophenol 1738-68-7, Glycine benzyl ester
                                                           1885-14-9, Phenyl
     chloroformate 2712-78-9, [Bis(trifluoroacetoxy)iodo]benzene
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Absolute stereochemistry.

IT

RN

CN

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L155 ANSWER 3 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
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     131:228660
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     Entered STN: 23 Sep 1999
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     Preparation of carbamoyloxymethyltetrahydroisoquinolinylalkanols as
TI
     central nervous system agents.
IN
     Choi, Yong Moon
PA
     SK Corp., USA
SO
     U.S., 14 pp.
     CODEN: USXXAM
DΥ
     Patent
     English
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     ICS C07D217-00; C07D217-16
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OS
     MARPAT 131:228660
GT
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$$\begin{array}{c|c} X^2 & R^4 & Y \\ \hline X^1 & NR^1 & NR^2R^3 \end{array}$$

TT

Ι

Title compds. (I; X1, X2 = H, alkyl, alkoxy, thioalkoxy, halo, OH, NO2, AΒ CF3; Y = O, S; R1 = H, alkyl, arylalkyl, CONHR'; R' = H, alkyl, arylalkyl, aryl; R2, R3 = H, alkyl, arylalkyl, cycloalkyl; R2R3N = 5-7 membered ring; R4 = H, alkyl), were prepared Thus, (S)-3-hydroxymethyl-1,2,3,4tetrahydroisoquinoline was PhCH2O2CCl and Na2CO3 in THF to give (S) - N-benzyloxycarbonyl-3-hydroxymethyl-1,2,3,4-tetrahydroisoquinoline.This in THF was treated with carbonyldiimidazole and then with aqueous NH3 to give (S)-N-benzyloxycarbonyl-3-carbamoyloxycarbonyl-1,2,3,4tetrahydroisoquinoline. The latter was hydrogenated in MeOH over Pd/C to give (S)-3-carbamoyloxymethyl-1,2,3,4-tetrahydroisoquinoline. The latter at 10 µM gave 97.7% inhibition of monoamine oxidase. carbamoyloxymethyltetrahydroisoquinolinylalkanol prepn central nervous ST system agent; isoquinolinylalkanol carbamoyloxymethyl prepn central nervous system agent; monoamine oxidase inhibitor carbamoyloxymethyltetrahydroisoquinolinylalkanol prepn; antidepressant carbamoyloxymethyltetrahydroisoquinolinylalkanol prepn IT Antidepressants Nervous system agents (preparation of carbamoyloxymethyltetrahydroisoguinolinylalkanols as central nervous system agents) 243858-56-2P 243858-57-3P 243858-58-4P IT 243858-59-5P 243858-61-9P 243858-62-0P 243858-69-7P 243858-70-0P 243858-68-6P 243858-65-3P 243858-72-2P **243858-74-4P** 243858-76-6P 243858-71-1P 243858-78-8P 243870-46-4P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of carbamoyloxymethyltetrahydroisoquinolinylalkanols as central nervous system agents)

nervous system agents)

103-71-9, Phenyl isocyanate, reactions 18881-17-9,

(S)-3-Hydroxymethyl-1,2,3,4-tetrahydroisoquinoline 41234-43-9, Ethyl
1,2,3,4-tetrahydroisoquinoline-3-carboxylate 59291-28-0
62855-02-1, (R)-3-Hydroxymethyl-1,2,3,4-tetrahydroisoquinoline
63006-93-9, 3-Hydroxymethyl-1,2,3,4-tetrahydroisoquinoline
243858-81-3 243858-83-5

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of carbamoyloxymethyltetrahydroisoquinolinylalkanols as central

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             THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
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RE
(1) Anon; GB 2266529 1993 HCAPLUS
(2) Anon; EP 564193 1993 HCAPLUS
(3) Anon; WO 9320099 1993 HCAPLUS
(4) Anon; WO 9413661 1994 HCAPLUS
(5) Anon; WO 9413664 1994 HCAPLUS
(6) Anon; WO 9617610 1994 HCAPLUS
(7) Anon; WO 9533727 1995 HCAPLUS
(8) Anon; WO 9616982 1996 HCAPLUS
(9) Blankley; US 5246943 1993 HCAPLUS
(10) Gafurov, M; Uzb Khim Zh 1988, V5, P15
(11) Gray, A; DE 1806900 HCAPLUS
(12) Kametani
(13) Kametani; 1968, V88(5), P573 HCAPLUS
(14) Renat; US 3449360 1969 HCAPLUS
(15) Richard; US 3308128 1967 HCAPLUS
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RN ·
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     3-Isoquinolinemethanol, 1,2,3,4-tetrahydro-, carbamate (ester), (3S)-
CN
           (CA INDEX NAME)
Absolute stereochemistry. Rotation (-).
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L155 ANSWER 4 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
     1992:151207 HCAPLUS
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DN
     Entered STN: 17 Apr 1992
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     Simple synthesis of N-(1-adamantyl)carbamates
TI
     Klimochkin, Yu. N.; Moiseev, I. K.
AU
     Kuibyshev. Politekh. Inst., Kuibyshev, USSR
CS
     Zhurnal Organicheskoi Khimii (1991), 27(8), 1795-6
so
     CODEN: ZORKAE; ISSN: 0514-7492
DT
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CC
     24-8 (Alicyclic Compounds)
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OS
GI
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NHCO<sub>2</sub>R
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C07D413-04

The title compds., e.g., I (R = Me, Et, CH2CH2OEt, CH2CH2NMe2) were prepared AB by the treatment of 1-adamantanol or its nitrate ester with ROH and KOCN in 37-64% yields. carbamoylation adamantanol; adamantylcarbamate ST 64-17-5, Ethanol, reactions 67-56-1, Methanol, reactions ΙT 108-01-0, 2-(Dimethylamino)ethanol 110-80-5, 2-Ethoxyethanol RL: RCT (Reactant); RACT (Reactant or reagent) (carbamoylation of adamantanol derivs. with, and potassium cyanate) 590-28-3, Potassium cyanate IT RL: RCT (Reactant); RACT (Reactant or reagent) (carbamoylation of adamantanol with, and alcs.) TT 15598-87-5 RL: RCT (Reactant); RACT (Reactant or reagent) (carbamoylation of, with methanol and potassium cyanate) 136860-51-0P 136860-59-8P IT 25192-03-4P 59987-81-4P 139537-50-1P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) 768-95-6, Tricyclo[3.3.1.13,7]decan-1-ol 32314-61-7 IT RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with potassium cyanate and alcs.) 108-01-0, 2-(Dimethylamino) ethanol ITRL: SPN (Synthetic preparation); PREP (Preparation) (carbamoylation of adamantanol derivs. with, and potassium cyanate) RN108-01-0 HCAPLUS Ethanol, 2-(dimethylamino)- (8CI, 9CI) (CA INDEX NAME) CN $Me_2N-CH_2-CH_2-OH$ L155 ANSWER 5 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN 1988:610794 HCAPLUS ANDN 109:210794 Entered STN: 10 Dec 1988 ED3-Pyrrolidinylthio-1-azabicyclo [3.2.0]hept-2-ene-2-carboxylic acid TT compounds, their preparation, pharmaceuticals containing them, their use against infections, and intermediates and their preparation Murata, Masayoshi; Tsutsumi, Hideo; Matsuda, Keiji; Hattori, Kohji; IN Nakajima, Takashi Fujisawa Pharmaceutical Co., Ltd., Japan PΑ Eur. Pat. Appl., 79 pp. so CODEN: EPXXDW  $\mathbf{DT}$ Patent English LA TC ICM C07D487-04 ICS A61K031-40; C07D417-04; C07D403-04; C07D405-04; C07D409-04;

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CO2R6

 $R^7$ 

MARPAT 109:210794

OS GI

The title compds. I [R1 = (protected) carboxy; R2 = (protected) hydroxyalkyl; R3 = H, alkyl; R4 = (un) substituted heterocyclyl; R5 = H, imino protective group] and their salts, useful as antimicrobials, were prepared Carbapemen III (R6 = CH2C6H4NO2-4, R7 = CO2CH2C6H4NO2-4), prepared in 5 steps from (2S,4R)-2-carbamoyl-4-methylsulfonyloxy-1-(4-nitrobenzyloxycarbonyl) pyrrolidine and Me2NCH(OMe)2, was hydrogenolyzed to give III (R5 = R7 = H) (IV). In in vitro testing, the min. inhibitory concentration of IV against Proteus vulgaris 49 was 0.05 μg/mL.

ST antimicrobial azabicycloheptenecarboxylate prepn; bactericide azabicycloheptenecarboxylate prepn; carbapenem antimicrobial prepn

III

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Bactericides, Disinfectants, and Antiseptics
IT
     Fungicides and Fungistats
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IT
    Lactams
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                                   117336-38-6P 117336-41-1P
     117336-36-4P
                    117336-37-5P
                                                   117336-68-2P
                    117336-58-0P
                                    117336-67-1P
     117336-51-3P
                                    117336-71-7P
                                                   117336-72-8P
                                                                  117336-73-9P
     117336-69-3P
                    117336-70-6P
                                                                  117339-81-8P
                                    117336-79-5P
                                                   117336-98-8P
                    117336-77-3P
     117336-74-0P
                    117355-23-4P
     117355-22-3P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, as intermediate for carbapenem antimicrobials)
                                     60-23-1, 2-Aminoethanethiol
     56-45-1, L-Serine, reactions
IT
                                                              70-23-5, Ethyl
     62-55-5, Thioacetamide
                              62-56-6, Thiourea, reactions
                     76-83-5, Trityl chloride
                                                            107-15-3,
                                                 100-79-8
     bromopyruvate
                                     107-21-1, 1,2-Ethanediol, reactions
     1,2-Ethanediamine, reactions
     109-80-8, 1,3-Propanedithiol
                                     115-08-2, Thioformamide
                                                               302-01-2,
                            507-09-5, Ethanethioic acid, reactions
                                                                       540-63-6,
     Hydrazine, reactions
     1,2-Ethanedithiol 590-28-3, Potassium cyanate
```

```
1189-71-5, Chlorosulfonyl isocyanate
                                                       4457-32-3,
     4-Nitrobenzyloxycarbonyl chloride 4637-24-5
                                                     4704-77-2,
                               5470-11-1, Hydroxylamine hydrochloride
     3-Bromo-1,2-propanediol
                                           22483-09-6
                                                        26628-22-8, Sodium
     6610-29-3, 4-Methylthiosemicarbazide
                                                               89226-13-1
            36016-40-7, O-(Mesitylenesulfonyl)hydroxylamine
     90822-24-5 96035-08-4 96035-09-5
                                       117333-90-1
                                 117334-67-5
                                                              117336-21-7
                                               117336-17-1
     117334-02-8
                  117334-08-4
     117336-30-8 117336-40-0 117336-42-2
                   117336-52-4
                                 117336-60-4 117336-65-9
     117336-45-5
                                               117336-86-4
                                                              117355-20-1
                   117336-78-4
                                 117336-80-8
     117336-77-3
                   117407-10-0
     117355-24-5
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, in synthesis of antimicrobial carbapenems)
IT
     117333-89-8P
     RL: RCT (Reactant); RACT (Reactant or reagent);
     SPN (Synthetic preparation); RACT (Reactant or reagent);
     PREP (Preparation)
        (preparation and reaction of, in synthesis of antimicrobial carbapenems)
RN
     117333-89-8 HCAPLUS
     1-Pyrrolidinecarboxylic acid, 2-(aminocarbonyl)-4-[(triphenylmethyl)thio]-
CN
      (4-nitrophenyl) methyl ester, (2S-cis) - (9CI) (CA INDEX NAME)
```

Absolute stereochemistry.

1987:130551 HCAPLUS

AN

L155 ANSWER 6 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN

```
DN
     106:130551
     Entered STN: 17 Apr 1987
ED
     Synthesis and reactivities of triisocyanatoantimony
ΤI
     Kijima, Ichiro; Wakeshima, Ikuko; Sasaki, Toru
ΑU
     Fac. Eng., Sci. Univ. Tokyo, Tokyo, 162, Japan
CS
     Nippon Kagaku Kaishi (1986), (12), 1754-57
SO
     CODEN: NKAKB8; ISSN: 0369-4577
     Journal
DT
LA
     Japanese
     78-5 (Inorganic Chemicals and Reactions)
CC
     CASREACT 106:130551
os
     Sb(NCO)3 was prepared by the reaction of SbCl3 with NaOCN in the
AB
     presence of several additives in benzene and THF. The reaction was
     accelerated remarkably by using THF as an additive in benzene to give
     Sb(NCO)3 in high yield. Sb(NCO)3 reacted with amines such as NHEt2,
     BuNH2, PhH2, and NH3 to afford only the corresponding triureidoantimony
     compds., but reacted with alcs. such as iso-PrOH, BuOH, sec- and tert-BuOH
     or PhOH to yield the corresponding carbamate and trialkoxo- or
     triphenoxyantimony compds. Sb(NCO)3 reacted also with
     2-diethylaminoethanol (HL) to give SbL3 and 2-diethylaminoethyl carbamate,
     together with isocyanuric acid. Sb(NCO)3 reacted with alcs. and PhOH to
     yield the corresponding substituted products, but the reaction with amines
     provided only the corresponding addition products.
```

```
antimony isocyanate prepn reactivity; isocyanate
ST
     antimony prepn reactivity; amine reaction antimony isocyanate;
     alc reaction antimony isocyanate; sodium
     cyanate reaction antimony chloride; stibine trichloro reaction
     sodium cyanate
     Alcohols, reactions
TΤ
     Amines, reactions
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with antimony triisocyanate in benzene)
     67-64-1, Acetone, uses and miscellaneous 68-12-2, uses and miscellaneous
TT
                                      75-52-5, Nitromethane, uses and
     75-05-8, uses and miscellaneous
     miscellaneous 109-99-9, uses and miscellaneous
                                                      123-91-1, uses
     and miscellaneous
                        141-78-6, Ethyl acetate, uses and miscellaneous
     RL: USES (Uses)
        (antimony trichloride reaction with sodium cyanate
        in benzene containing, antimony triisocyanate formation in
        relation to)
TΤ
     108-20-3, Diisopropyl ether
                                   680-31-9, reactions
                                                         4067-16-7,
                            25322-68-3, Polyethylene glycol
                                                              26027-38-3
     Pentaethylenehexamine
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (antimony trichloride reaction with sodium cyanate
        in benzene containing, antimony triisocyanate formation in
        relation to)
     17455-13-9
TТ
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (antimony trichloride reaction with sodium cyanate
        in benzene or THF containing, antimony triisocyanate formation in
        relation to)
TT
     86893-88-1P, Antimony triisocyanate
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (preparation and reactivity of)
                                  2155-74-0P, Antimony tributoxide
     592-35-8P, Butyl carbamate
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with Bu
        alc. in benzene)
IT
     107320-93-4P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with
        ammonia in benzene)
TT
     107320-92-3P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with
        aniline in benzene)
     107320-91-2P
TΤ
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with
        butylamine)
TT
     107320-90-1P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with
        diethylamine in benzene)
IT
     108-80-5P, Isocyanuric acid 60743-30-8P, 2-Diethylaminoethyl
     carbamate 107295-94-3P, Tris(2-diethylaminoethoxo)antimony
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with
        diethylaminoethanol)
IT
     1746-77-6P, Isopropyl carbamate
                                       18770-47-3P, Antimony triisopropoxide
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, by reaction of antimony triisocyanate with iso-Pr
        alc. in benzene)
IΤ
     622-46-8P, Phenylcarbamate
                                  16484-27-8P, Antimony triphenoxide
     RL: SPN (Synthetic preparation); PREP (Preparation)
```

(preparation of, by reaction of antimony triisocyanate with phenol

kumar - 10 / 680979 in benzene) 93913-73-6P, Antimony tri-sec-butoxide 2114-15-0P, sec-Butyl carbamate IT RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by reaction of antimony triisocyanate with sec-Bu alc. in benzene) 10433-03-1P, Antimony tri-tert-butoxide 4248-19-5P, tert-Butyl carbamate IT RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, by reaction of antimony triisocyanate with tert-Bu alc. in benzene) 917-61-3, Sodium cyanate IT RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with antimony trichloride in benzene, effect of additives on antimony triisocyanate formation in) 62-53-3, Aniline, reactions 67-63-0, Isopropyl alcohol, reactions IT 71-36-3, reactions 75-65-0, tert-Butyl alcohol, reactions sec-Butyl alcohol 100-37-8, 2-(Diethylamino)ethanol 108-95-2, 109-89-7, reactions 7664-41-7, 109-73-9, reactions reactions Ammonia, reactions RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with antimony triisocyanate in benzene) 10025-91-9, Antimony chloride (SbCl3) IT RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with sodium cyanate in benzene, effect of additives on antimony triisocyanate formation in) 75-05-8, uses and miscellaneous ΙŤ RL: RCT (Reactant); RACT (Reactant or reagent) (antimony trichloride reaction with sodium cyanate in benzene containing, antimony triisocyanate formation in relation to)

### $H_3C-C=N$

RN

CN

75-05-8 HCAPLUS

L155 ANSWER 7 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN ΑN 1984:571174 HCAPLUS 101:171174 DN Entered STN: 10 Nov 1984 ED 2-Aryl-5,5-dimethyl-1,2,4-triazolidin-3-one derivatives TТ Schantl, J.; Hebeisen, P. ΑU Inst. Org. Pharm. Chem., Univ. Innsbruck, Innsbruck, A-6020, Austria CS Scientia Pharmaceutica (1983), 51(4), 379-90 SO CODEN: SCPHA4; ISSN: 0036-8709 DTJournal LA German 28-10 (Heterocyclic Compounds (More Than One Hetero Atom)) CC GΙ

Acetonitrile (8CI, 9CI) (CA INDEX NAME)

AB RnC6H5-nNHN:CMe2 [Rn = H, 4-Cl, 3,4-Cl2, 4-Me(CH2)50, 4-02N] reacted with KZCN (Z = 0, S) in AcOH to give the corresponding triazolidinones I (Z = 0) or -thiones I (Z = S). Although I (Z = S) have antiinflammatory and

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analgesic properties I (Z = O) had no noteworthy activity.
    RnC6H5-nN:NCMe2N:C:Z, the acyclic oxidation products of I, can be used for
     further syntheses. H2NCN was added to 4-ClC6H4NHN: CMe2. HCl to give
     iminotriazolidine II which on oxidative ring cleavage gave
     4-C1C6H4N:NCMe2NHCN.
     acetone hydrazone cyclization cyanate thiocyanate;
ST
     triazolidinone prepn oxidn; triazolidinethione prepn oxidn; cyanamide addn
     hydrazone
TT
     Analgesics
     Inflammation inhibitors and Antiarthritics
        (triazolidinethiones)
                               91027-26-8
                                            91027-27-9
                                                          91027-28-0
                28359-16-2
IT
     18440-33-0
                 91027-30-4
     91027-29-1
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (hydrolysis or ammonolysis of)
                           5877-04-3
                                        28359-15-1
     103-02-6 1200-11-9
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (phosgenation and ammonolysis of, or reaction with potassium
        cyanate or potassium thiocyanate)
     91027-31-5P
IT
                  91027-32-6P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and hydration of)
     18440-37-4P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and hydrogenolysis of)
                                 59395-36-7P
                                                59395-39-0P
                                                              72731-37-4P
                   39263-68-8P
ΤT
     24648-29-1P
     72731-38-5P
                   73150-88-6P
                                 91027-23-5P
                                                91027-24-6P
                                                              91027-25-7P
     91027-37-1P
                   91027-38-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and oxidation of)
                   91027-19-9P
                                91027-20-2P
IT
     91027-18-8P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reaction of, with potassium cyanate or
        thiocyanate)
IT
     91027-36-0P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
                                  124-02-7
                                               91027-33-7
                                                            91027-34-8
TT
               108-00-9 108-01-0
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with (arylazo) isocyanatopropane)
                 19763-90-7
                              91027-17-7
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with acetone)
     420-04-2
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with acetone (chlorophenyl) hydrazone)
IT
     333-20-0 590-28-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with acetone phenylhydrazones)
     91027-35-9
TT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with guanidine)
                               91027-21-3
                                             91027-22-4
                  57883-13-3
TT
     53670-11-4
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with potassium cyanate or
        thiocyanate)
IT
     91027-36-0P
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of)
```

RN

91027-36-0 HCAPLUS

CN Carbamic acid, [1-[(4-chlorophenyl)azo]-1-methylethyl]-, 2-(dimethylamino)ethyl ester, monohydrochloride (9CI) (CA INDEX NAME)

$$N = N - C - Me$$

$$M = N - C - Me$$

$$M = Me$$

### HCl

HU 194876

В

19880328

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L155 ANSWER 8 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
     1984:209515 HCAPLUS
AN
     100:209515
DN
     Entered STN: 23 Jun 1984
ED
     1-Sulfo-2-azetidinone derivatives
TT
     Kishimoto, Shoji; Matsuo, Taisuke; Ochiai, Michihiko
TN
     Takeda Chemical Industries, Ltd. , Japan
PA
     Eur. Pat. Appl., 186 pp.
SO
     CODEN: EPXXDW
DT
     Patent
     English
LA
     C07D205-08; C07D417-12; C07D417-14; C07D401-06; C07D413-14; C07D277-40;
IC
     C07D277-42; A61K031-365; A61K031-42; A61K031-425
     26-5 (Biomolecules and Their Synthetic Analogs)
CC
     Section cross-reference(s): 10, 63
FAN.CNT 6
     PATENT NO.
                                DATE
                                             APPLICATION NO.
                                                                    DATE
                         KIND
                                             ______
     .
_______
                         _ _ _ _
                                -----
                                             EP 1983-104061
                                                                    19830426 <--
PI
     EP 93376
                          A2
                                19831109
     EP 93376
                          Α3
                                19861015
                                19900321
                          В1
     EP 93376 /
                          B2
                                19990407
     EP 93376
         R: AT, BE, CH, DE, FR, IT, LI, LU, NL, SE
                                                                    19820430 <--
                                             JP 1982-73728
                                19831104
     JP 58189176
                          A2
                                19880708
                          B4
     JP 63034155
                                             JP 1982-93463
                                                                    19820531 <--
                          A2
                                19831207
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                          B4
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                                             AU- 1983-13445
                                                                    19830412 <--
                         · A1
                                19831103
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     AU 564150
                          B2
                                19870806
                                                                     19830419 <--
                          Α
                                19831228
                                             ZA 1983-2742
     ZA 8302742
                                             GB 1983-10520
                                                                     19830419 <--
                          A1
                                19840215
     GB 2124207
                          B2
                                19861008
     GB 2124207
                                             AT 1983-104061
                                                                     19830426 <--
                          E
                                 19900415
     AT 51223
                                             DK 1983-1889
                                                                     19830428 <--
     DK 8301889
                          Α
                                 19831031
     DK 161832
                          В
                                 19910819
                          С
                                 19920120
     DK 161832
                          Α
                                 19831031
                                             FI 1983-1457
                                                                     19830428 <--
     FI 8301457
                                             SU 1983-3590552
                                                                    19830428 <--
     SU 1480763
                          A3
                                 19890515
                                             NO 1983-1514
                                                                     19830429 <--
     NO 8301514
                          Α
                                 19831031
     NO 160581
                          В
                                 19890123
                          С
                                 19890503
     NO 160581
                          0
                                 19840328
                                             HU 1983-1486
                                                                    19830429 <--
     HU 30672
```

$$R^{1}$$
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{2$ 

CASREACT 100:209515

OS GI A61K031-425

The title compds. I [R = COR3, (CH2) nR4, N-containing heterocyclyl; R1 = AB (un)acylated or (un)protected amino group; R2 = H, MeO; R3 = (un)protected or (un) substituted NH2, (un) protected OH; R4 = H, halo, NHCONH2, NHCONHSO3H, CONH2, O2CNH2, O2CNHSO3H, alkylsulfonyloxy, pyridinio, alkoxy, alkylsulfinyl, -sulfonyl, haloalkylcarbonyloxy, OH, alkoxycarbonyl, acyloxy, alkoxyiminoalkyl, alkylcarbonyl, acylamine; n = 1-3] or their salts or esters, with improved antimicrobial and  $\beta$ -lactamase inhibitory activity, were prepared Thus, sulfonating (3S,4S)-cis-3benzyloxycarboxamido-4-carbamoyloxymethyl-2-azetidinone in dioxane with SO3-pyridine complex at room temperature 14 h and converting the product to the Na salt gave 64% Na (3S,4S)-cis-3-benzyloxycarboxamido-4carbamoyloxymethyl-2-acetidinone-1-sulfonate. Hydrogenolysis of the latter removed the amino protective group and the product was acylated with a substituted acetyl chloride-HCl and then hydrolyzed with MeNHCO2Na to give 76% acetamidoazetidinonesulfonate (3S,4S)-cis-(Z)-II (R5 =

```
This was deprotected by room temperature
     4-02NC6H4CH2, R6 = Na).
hydrogenolysis
     to give 61\% (3S,4S)-cis-(Z)-II (R5 = R6 = H) (III). III had a min.
     inhibitory concentration of 0.05 \mu g/mL against Enterobacter cloacae IFO 12937
     and Klebsiella pneumoniae TN 1711 and 1.56 µg/mL against Pseudomonas
     aeruginosa GN 3407.
     bactericide azetidinonesulfonate prepn; sulfoazetidinone bactericide prepn
ST
     Bactericides, Disinfectants, and Antiseptics
IT
        (sulfoazetidinone derivs.)
IT
     83175-92-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (Wittig methylenation of)
                84186-87-8
                            84208-30-0
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of aminoazetidinone derivative)
IT
     84208-37-7
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of aminoazetidinone derivs.)
IT
     65243-22-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of aminoazetidinones)
     24424-99-5
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of aminoazetidione derivative)
     76903-12-3
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of glycine derivative)
     41295-64-1
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of nitrobenzyl alc.)
     22818-40-2
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation of, with chromonecarbonyl chloride derivative)
     16869-24-2
                  23877-12-5
TT
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (alkylation by, of (hydroxyimino)acetate)
     64485-82-1
TT
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (alkylation of, by bromopropionate ester)
IT
     1668-10-6
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (amidation by, of azetidinonecarboxylic acid)
IT
     79656-47-6
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (amidation of, by aminoazetidinone derivative)
     84208-16-2
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (amidation of, by glycine amide)
     90121-74-7 90121-75-8 90121-76-9
      90121-77-0 90121-78-1 90121-79-2
      90121-80-5 90192-20-4 90192-21-5
      90192-22-6 90192-23-7 90192-24-8
      90242-01-6 90242-02-7
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
      study, unclassified); BIOL (Biological study)
         (bactericidal activity of)
      61964-78-1
IT
      RL: RCT (Reactant); RACT (Reactant or reagent)
         (butoxycarbonylation of and conversion to tartrate salt)
      5470-11-1
 IT
      RL: RCT (Reactant); RACT (Reactant or reagent)
         (cyclization of, with (acetyloxopropyl)azetidinone derivative)
TT
      371-62-0
```

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RL: RCT (Reactant); RACT (Reactant or reagent)
        (cyclization of, with dimethoxybenzylamine and phthaloylglycine
        chloride)
IT
     6780-38-7
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (cyclization of, with fluoroethanol and dimethoxybenzylamine)
IT
     20781-20-8
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (cyclization of, with fluoroethanol and phthaloylglycine chloride)
IT
     83422-65-5
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (cyclization of, with hydroxylamine)
     62-56-6, reactions
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (cyclization of, with nitrobenzyl (hydroxyimino)chloroacetoacetate)
     90121-41-8
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (deprotection and acylation of, with acetyl chloride derivative)
TΤ
     120-78-5
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (esterification by, of (methoxyimino) acetic acid derivative)
TT
     619-73-8
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (esterification of, with chloroacetoacetyl chloride)
                  76134-88-8
                               90121-52-1
                                             90121-53-2
     76134-87-7
TT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (hydrogenation of)
     84209-04-1
TΤ
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (hydrogenolysis of)
IT
     84186-82-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (hydrolysis of)
                                 90121-66-7P 90192-19-1P
IT
     88792-29-4P
                   90121-58-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and acylation of, with acetyl chloride derivative)
     87638-04-8P 88852-08-8P 90121-38-3P
IT
     90192-08-8P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL
     (Biological study); PREP (Preparation)
        (preparation and antibacterial activity of)
IT
     90121-48-5P 90192-25-9P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation)
        (preparation and bactericidal activity of)
IT
     90192-05-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and benzyloxycarbonylation of)
     90121-64-5P
TT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and blocking of, carbobenzoxy chloride)
     90192-26-0P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and conversion of, to sodium salt)
IT
     90121-84-9P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and conversion to sodium salt)
     86832-68-0P
IT
     RL: PRP (Properties); SPN (Synthetic preparation); PREP
```

```
(Preparation)
        (preparation and crystal structure of)
IT
     90121-72-5P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and cyclization of, with thiourea)
                              89605-11-8P 90121-17-8P
     86299-59-4P 86334-64-7P
IT
     90121-65-6P 90121-70-3P
     RL: RCT (Reactant); SPN (Synthetic preparation);
     PREP (Preparation); RACT (Reactant or reagent)
        (preparation and debenzylation of)
                   86791-57-3P 89604-55-7P
IT
     86299-42-5P
                   90121-45-2P 90121-54-3P
     90121-42-9P
     90192-10-2P 90192-12-4P
     RL: RCT (Reactant); SPN (Synthetic preparation);
     PREP (Preparation); RACT (Reactant or reagent)
        (preparation and deprotection of)
IT
     90121-59-8P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and desilylation of)
     86299-57-2P
TΤ
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and ester cleavage of)
     74440-02-1P 86299-47-0P
TΤ
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and esterification of, with benzothiazolyl disulfide)
IT
     86299-58-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and esterification of, with dithiobisbenzothiazole)
                   90121-63-4P
TΤ
     90121-57-6P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and hydrazinolysis of)
IT
     90121-24-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and hydrogenation of)
IT
     88124-54-3P
                   90121-26-9P 90121-37-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
    (Preparation); RACT (Reactant or reagent)
        (preparation and hydrogenolysis of)
IT
     90121-47-4P 90121-61-2P 90121-69-0P
     90121-83-8P 90192-16-8P
     RL: RCT (Reactant); SPN (Synthetic preparation);
     PREP (Preparation); RACT (Reactant or reagent)
        (preparation and hydrolysis of)
IT
     122645-63-0P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and mesylation of)
     90121-28-1P 90121-82-7P
                               90192-06-6P
TT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and oxidation of)
     90121-71-4P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
         (preparation and oximation of)
     90121-56-5P
TΤ
     RL: SPN (Synthetic preparation); PREP (Preparation)
```

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(preparation and protection of, with silyl chloride derivative)
                   89605-09-4P
IT
     86299-52-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reaction of, with aminoazetidinone derivative)
     86334-63-6P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and reaction of, with chlorosulfonyl isocyanate)
IT
     90121-39-4P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and reaction of, with potassium cyanate)
     90121-73-6P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reaction of, with tert-Bu bromoacetate)
     61964-79-2P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reduction of)
     86299-46-9P
TT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and saponification of)
     84209-18-7P 90121-36-1P 90121-60-1P
IT
     90121-68-9P 90121-81-6P 90192-15-7P
     RL: RCT (Reactant); SPN (Synthetic preparation);
     PREP (Preparation); RACT (Reactant or reagent)
        (preparation and sulfonation of)
IT
     89604-92-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and N-acylation by, of aminoazetidione derivative)
     84209-05-2P 88852-06-6P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation);
     PREP (Preparation); RACT (Reactant or reagent)
        (preparation and N-acylation of, by acetyl chloride derivative)
IT
     90121-27-0P
                   90121-46-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and N-acylation of, with acetyl chloride derivative)
     69284-11-3P 82935-61-3P 84187-15-5P
IT
     90121-15-6P
                   90121-18-9P 90121-19-0P
                   90121-25-8P 90121-29-2P
     90121-20-3P
     90121-30-5P 90121-31-6P
                               90121-32-7P
     90121-33-8P 90121-40-7P 90121-43-0P
     90121-49-6P 90121-51-0P 90121-55-4P
                   90192-07-7P 90192-11-3P
     90121-62-3P
     90192-13-5P 90192-17-9P 90244-80-7P
     122645-64-1P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
IT
     86334-65-8P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation, sulfonation, and hydrogenolysis of)
TΤ
     18162-48-6
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (protection by, of azetidinone derivs.)
IT
     590-28-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with (aminomethyl)azetidinone derivative)
```

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2605-67-6
IT
     1189-71-5
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with (hydroxymethyl)azetidinone derivative)
     86299~56-1
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with bromoacetate)
IT
     83175-92-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with chlorosulfonyl isocyanate and
        debenzylation of)
     5292-43-3
IT
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with hydroxyiminoacetate derivative)
     90121-16-7 90121-50-9
TT
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with potassium cyanate)
TΤ
     90121-44-1
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with pyridine)
                90192-18-0
IT
     84187-90-6
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reduction of)
IT
     9073-60-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (sulfoazetidinones as inhibitors of)
IT
     90121-34-9
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (sulfonation and hydrolysis of)
IT
     86299-43-6
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (N-acylation of, by benzothiazolyl thioacetate derivative)
IT
     88792-25-0
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (O-methylation of)
IT
     83175-92-2
    RL: RCT (Reactant); RACT (Reactant or reagent);
    PREP (Preparation); RACT (Reactant or reagent);
    PREP (Preparation)
        (Wittig methylenation of)
RN
     83175-92-2 HCAPLUS
     Carbamic acid, [1-[(2,4-dimethoxyphenyl)methyl]-2-(hydroxymethyl)-4-oxo-3-
CN
     azetidinyl]-, phenylmethyl ester, cis- (9CI) (CA INDEX NAME)
```

Relative stereochemistry.

```
L155 ANSWER 9 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN AN 1979:419926 HCAPLUS
DN 91:19926
ED Entered STN: 12 May 1984
TI N-Substituted carbamates
IN Chung, Rack H.
PA BASF Wyandotte Corp., USA
```

```
SO
    U.S., 5 pp.
    CODEN: USXXAM
DT
    Patent
LA
    English
IC
    C07C125-04
NCL
    260465000D
    23-20 (Aliphatic Compounds)
CC
    Section cross-reference(s): 25
FAN.CNT 1
                                        APPLICATION NO.
                                                               DATE
                       KIND DATE
    PATENT NO.
    _____
                                         -----
                                                                ------
                       ____
                              _____
    US 4147716
                              19790403 US 1978-912461
                                                               19780605 <--
                        A
                              19780605 <--
PRAI US 1978-912461
CLASS
               CLASS PATENT FAMILY CLASSIFICATION CODES
PATENT NO.
 ______
                      C07C125-04
               IC
US 4147716
               NCT.
                      260465000D
    The title compds. were prepared by the reaction of RX (R = alkyl, alkenyl,
AB
    aralkyl, aralkenyl; X = halo) with alkali cyanates and
    monohydric and polyhydric alcs. (not aromatic) at 65-100° in
    sulfolane. Thus, KOCN in sulfolane was heated to 90°,
    EtBr was added in 1.5 h, PhN(CH2CH2CN)CH2CH2OH in sulfolane was added, and
    the mixture was heated 3 h at 90° and worked up to give
    EtNHCO2CH2CH2N (CH2CH2CN) Ph.
    alkylcarbamate anilinoethyl; alc alkali cyanate haloalkane
ST
    alkylcarbamate
    63216-95-5P 70489-11-1P
TT
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (preparation of)
IT
    92-64-8
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (reaction with alkyl halides and potassium cyanate,
       carbamate esters from)
IT
     590-28-3
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (reaction with anilinoethanol derivative and alkali halides, carbamate
       esters, from)
IT
              75-00-3
                       106-94-5
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (reaction with potassium cyanate and anilinoethanol
       derivative, carbamate ester from)
IT
     63216-95-5P
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (preparation of)
RN
     63216-95-5 HCAPLUS
    Carbamic acid, ethyl-, 2-[(2-cyanoethyl)phenylamino]ethyl ester (9CI) (CA
CN
     INDEX NAME)
     0
                  Ph
{\tt EtNH-C-O-CH_2-CH_2-N-CH_2-CH_2-CN}
L155 ANSWER 10 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
AΝ
    1978:170147 HCAPLUS
DN
    88:170147
ED
    Entered STN: 12 May 1984
    1-Methyl-2-(carbamyloxymethyl)-5-nitroimidazole
TI
   FARCHEMIA di Martino Finotto e C. S.a.S., Italy
PA
```

Belg., 9 pp.

```
CODEN: BEXXAL
    Patent
\mathbf{DT}
    French
LA
    C07D
IC
    28-10 (Heterocyclic Compounds (More Than One Hetero Atom))
CC
FAN.CNT 1
                      KIND DATE
                                       APPLICATION NO.
                                                              DATE
    PATENT NO.
                       ----
                             _____
    _____
                      A1 19770916 BE 1977-55955
A1 19781201 FR 1977-16498
                                                              19770531 <--
    BE 855188
PΙ
                                                              19770531 <--
    FR 2389609
                      B1 19800711
    FR 2389609
                                      NL 1977-6061
                       Α
                            19781107
                                                              19770602 <--
    NL 7706061
                      В
    NL 175298 .
                            19840528
                      C
                            19841016
    NL 175298
                            19770503 <--
PRAI IT 1977-23102
CLASS
              CLASS PATENT FAMILY CLASSIFICATION CODES
PATENT NO.
 BE 855188 IC
                     C07D
    The title compound was obtained in 93% yield by treating
AB
    1-methyl-2-hydroxymethyl-5-nitroimidazole with 1-carbamoylimidazole.
    carbamoyloxymethylimidazole; imidazole carbamoyloxymethyl
ST
    7681-76-7P
IT
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (preparation of)
    936-05-0
IT
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (reaction of, with carbamoylimidazole)
IT
     2578-41-8 66339-05-7
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (reaction of, with hydroxymethylimidazole derivative)
IT
    616-47-7
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (reaction of, with potassium cyanate and
       hydroxymethylimidazole derivative)
IT
     7681-76-7P
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (preparation of)
RN
     7681-76-7 HCAPLUS
     1H-Imidazole-2-methanol, 1-methyl-5-nitro-, carbamate (ester) (9CI) (CA
CN
     INDEX NAME)
```

$$O_2N \xrightarrow{N} CH_2 - O - C - NH_2$$

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L155 ANSWER 11 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
    1975:458900 HCAPLUS
AN
     83:58900
DN
    Entered STN: 12 May 1984
ED
     2,3-Benzoxazepine derivatives
TI
     Pifferi, Giorgio; Omodei-Sale, Amedeo; Consonni, Pietro
IN
     Gruppo Lepetit S.p.A., Italy
PΑ
SO
     Can., 15 pp.
     CODEN: CAXXA4
DT
     Patent
LA
     English
```

```
28-24 (Heterocyclic Compounds (More Than One Hetero Atom))
CC
FAN.CNT 1
                       KIND
                               DATE
                                          APPLICATION NO.
                                                                 DATE
    PATENT NO.
                                                                 _____
                        ____
                                          -----
                                          CA 1972-137695 19720321 <--
    CA 959054
                        A1
                               19741210
PT
                               19720321 <--
PRAI CA 1972-137695
CLASS
 PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
 _____
CA 959054
    For diagram(s), see printed CA Issue.
    Antiinflammatory (no data) benzoxazepines I (R = alkyl, alkenyl,
     carbamoyl, acyl, alkoxycarbonyl) (26 compds.) were prepared in 39-91% yield
    by a) methylation of I (R = H), b) alkylation of I (R = H) with alkyl and
     alkenyl halides, c) acylation of I (R = H) with acyl halides, d) treatment
     of I (R = H) with isocyanates and isothiocyanates, and
     e) treatment of I (R = COCl) with alkylamines, morpholine, pyrrolidines,
     and piperazines. The cycloaddn. of 2-BrCH2C6H4CH2CH2Br with KONHCO2Et
     gave 79% I (R = CO2Et) which was hydrolyzed-decarboxylated to 74% I (R =
     H). I (R = COCl) was obtained in 81.5\% yield by treating I (R = H) with
     COC12. Also I were central nervous system depressants.
     benzoxazepine central depressant antiinflammatory; carbamoylbenzoxazepine;
ST
     alkylation benzoxazepine; acylation benzoxazepine
     Inflammation inhibitors
IT
        (benzoxazepines as)
     Nervous system
TT
        (depressant for central, benoxazepines as)
     Cycloaddition reaction
IT
        (of (bromomethyl) phenethyl bromide with hydroxyurethane, benzoxazepine
        by)
     Acylation
IT
     Alkylation
        (of benzoxazepine)
     79-04-9 79-44-7 83-01-2 88-10-8 590-21-6
                                                        3350-78-5 4521-61-3
TT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (acylation by, of tetrahydrobenzoxazepine)
     78-77-3 106-95-6
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (alkylation by, of tetrahydrobenzoxazepine)
IT
     56190-13-7
     RL: PROC (Process)
        (cycloaddn. of, with (bromomethyl)phenethyl bromide)
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and hydrolysis-decarboxylation of)
     38090-29-8P
TT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
         (preparation and reaction of)
     35040-43-8P 35040-44-9P 35040-45-0P 35040-46-1P
IT
     35040-48-3P 35040-49-4P 35040-50-7P 35040-51-8P
                                                             35040-87-0P
     35040-88-1P 35040-89-2P 38090-25-4P 38090-28-7P
     38090-30-1P 38090-36-7P 38090-37-8P 38090-38-9P 38090-40-3P 38090-41-4P 38090-42-5P 38090-43-6P 38090-45-8P 38090-46-9P 38090-47-0P 38090-48-1P
                                                             38090-39-0P
                                                             38090-44-7P
     RL: SPN (Synthetic preparation); PREP (Preparation)
         (preparation of)
IT
     556-61-6
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (reaction of, with (hydroxyethyl)tetrahydrobenzoxazepine)
IT
     38256-56-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
```

```
(reaction of, with hydroxyurethane)
     103-71-9 624-83-9
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with tetrahydrobenzoxazepine)
             100-36-7 109-01-3 110-91-8 123-75-1
                                                         124-02-7 124-40-3,
TТ
     reactions 30381-48-7
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with tetrahydrobenzoxazepinecarbonyl chloride)
IT
     917-61-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with tetrahydrobenzoxazepineethanol)
     75-21-8, reactions
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (with tetrahydrobenzoxazepine)
IT
     35040-43-8P
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of)
     35040-43-8 HCAPLUS
RN
     2,3-Benzoxazepine-3(1H)-ethanol, 4,5-dihydro- (9CI) (CA INDEX NAME)
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CN

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L155 ANSWER 12 OF 12 HCAPLUS COPYRIGHT 2004 ACS on STN
    1975:125095 HCAPLUS
AN
    82:125095
DN
    Entered STN: 12 May 1984
ED
    2-Alkylaminobenzophenones
TI
    Welstead, William J., Jr.; Stauffer, Harold F., Jr.
IN
    A. H. Robins Co., Inc.
PA
    U.S., 6 pp.
SO
    CODEN: USXXAM
DT
    Patent
    English
T.A
IC
    C07D
NCL 260482000C
    25-16 (Noncondensed Aromatic Compounds)
    Section cross-reference(s): 1
FAN.CNT 1
                       KIND
                                         APPLICATION NO.
                                                               DATE
    PATENT NO.
                              DATE
                                         _____
     _____
                       ____
                              _____
                                                               19720920 <--
                                       US 1972-290568
                              19741105
    US 3846477
                       Α
                              19720920 <--
PRAI US 1972-290568
                CLASS PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
                      _ _ _ _
 _____
                IC
                      C07D
 US 3846477
                      260482000C
                NCL
     For diagram(s), see printed CA Issue.
GI
     5-Chloro-2-(tosylamido)benzophenone was treated with substituted alkyl
     halides and NaH to give the aminobenzophenones (I, R = H, CH2OH).
     Similarly prepared were the following II (n and R given): 1, H; 2, Me.
     N-methylation and N-acylation of the I gave 5,2-
     Cl[HOCH2CH(OH)CH2NMe]C6H3COPh and 5,2-Cl[HO(CH2)2N(CO2Et)]C6H3COPh which
     demonstrated tranquilizer activity.
     benzophenone hydroxyalkylamino tranquilizer; tranquilizer
ST
```

```
hydroxyalkylaminobenzophenone
IT
     Tranquilizers
        ([(hydroxyalkyl)amino]benzophenones)
IT
     541-41-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (N-acylation of [(hydroxyethyl)amino]benzophenone derivative by)
     79-44-7 917-61-3
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (O-carbamoylation of [(hydroxyethyl)amino]benzophenone derivative by)
     54524-10-6P 54524-12-8P
TΤ
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and N-methylation of, by formic acid-formaldehyde)
     33108-34-8P
ΤТ
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation and O-carbamoylation of)
IT
     54524-08-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reactions of)
     54524-13-9P 54524-15-1P
ΙT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and tranquilizer activity of)
                                54524-14-0P 54524-16-2P
                   54524-11-7P
     54524-09-3P
TΤ
     54524-17-3P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
                          13999-24-1
                                      51337-32-7
TT
               540-51-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with (tosylamido)benzophenone derivative)
     4873-59-0
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with alkyl halides)
ΙT
     917-61-3
     RL: SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (O-carbamoylation of [(hydroxyethyl)amino]benzophenone derivative by)
     917-61-3 HCAPLUS
RN
     Cyanic acid, sodium salt (8CI, 9CI) (CA INDEX NAME)
CN
HO-C=N
  Na
=> d l157 all hitstr tot
L157 ANSWER 1 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
     2004:182532 HCAPLUS
AN
     140:235607
DN
ED
     Entered STN: 05 Mar 2004
     Preparation of 4-benzoylpiperidine derivatives for treatment of psychosis
ΤI
     and cognition disorders
     Choi, Yong-moon; Kim, Yong-kil; Yoo, Jin-uk; Paek, Eun-ah; Park,
IN
     Chun-eung; Seo, Sung-yong; Chung, Coo-min; Heo, Joon
PA
     SK Corp., USA
     U.S. Pat. Appl. Publ., 30 pp.
SO
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CODEN: USXXCO
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     Patent
    English
LA
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     ICM A61K031-4709
     ICS A61K031-4545; A61K031-453; C07D041-02; C07D049-02
     514314000; 514317000; 514318000; 514326000; 546176000; 546194000;
NCL
     546225000; 546207000
     27-16 (Heterocyclic Compounds (One Hetero Atom))
CC
    Section cross-reference(s): 1
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                         A1
                               20040304
                                           US 2002-228869
                                                                  20020826
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    US 2004044033
                         B2
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            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
            PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
            TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG,
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                               20020826
PRAI US 2002-228869
                         Α
CLASS
 PATENT NO.
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 US 2004044033
                ICM
                       A61K031-4709
                       A61K031-4545; A61K031-453; C07D041-02; C07D049-02
                ICS
                NCL
                       514314000; 514317000; 514318000; 514326000; 546176000;
                       546194000; 546225000; 546207000
    MARPAT 140:235607
os
GI
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The title compds. I [wherein n = 0-2; A = thienyl, naphthyl, pyridyl, quinolyl, or (un)substituted Ph; X = 0-carbamoyl, alkoxy, imidazolyl, triazolyl, tetrazolyl, or carbonate; Y = H, halo, alkyl, or alkoxy] or racemic or enantiomerically enriched isomers, or pharmaceutically acceptable salts thereof are prepared For example, 4-(4-

II

ŧ

fluorobenzoyl)piperidine was reacted with (S)-4-isopropylstyrene oxide in isopropanol, followed by the addition of MsCl, Et3N, and MeOH to give II. II showed ED50 of 0.13 mg/kg as an antipsychotic agent in rat. I are useful for the treatment of central nervous system diseases in a mammal, in particular psychosis and cognition disorders. benzoyl piperidine treatment psychosis cognition disorder prepn Mental disorder

(cognitive; preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders)

IT Cognition

ST

IT

IT

IT

(disorder; preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders)

IT Antipsychotics

(preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders)

IT Mental disorder

(psychosis; preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders)

666858-06-6P 666858-07-7P 666858-08-8P 666858-12-4P 666858-13-5P 666858-09-9P 666858-10-2P 666858-11-3P 666858-16-8P 666858-17-9P 666858-18-0P 666858-14-6P 666858-15-7P 666858-21-5P 666858-22-6P 666858-23-7P 666858-19-1P 666858-20-4P 666858-25-9P 666858-26-0P 666858-27-1P 666858-28-2P 666858-24-8P 666858-29-3P 666858-30-6P 666858-31-7P 666858-32-8P 666858-33-9P 666858-35-1P 666858-36-2P 666858-38-4P 666858-39-5P 666858-34-0P 666858-41-9P 666858-42-0P 666858-43-1P 666858-44-2P 666858-40-8P 666858-45-3P 666858-46-4P 666858-47-5P 666858-48-6P 666858-49-7P 666858-50-0P 666858-51-1P 666858-52-2P 666858-53-3P 666858-54-4P 666858-59-9P 666858-55-5P 666858-56-6P 666858-57-7P 666858-58-8P 666858-60-2P 666858-61-3P 666858-62-4P 666858-63-5P 666858-64-6P 666858-69-1P 666858-65-7P 666858-66-8P 666858-67-9P 666858-68-0P 666858-75-9P 666858-77-1P 666858-70-4P 666858-71-5P 666858-73-7P 666858-85-1P 666858-87-3P 666858-79-3P 666858-81-7P 666858-83-9P 666858-89-5P 666858-91-9P 666858-93-1P 666858-95-3P 666858-97-5P 666859-03-6P 666859-05-8P 666859-07-0P 666858-99-7P 666859-01-4P 666859-13-8P 666859-15-0P 666859-17-2P 666859-09-2P 666859-11-6P 666859-21-8P 666859-23-0P 666859-25-2P 666859-27-4P 666859-19-4P 666859-29-6P 666859-31-0P 666859-33-2P 666859-35-4P 666859-37-6P 666859-39-8P 666859-41-2P 666859-45-6P 666859-47-8P 666859-49-0P 666859-51-4P 666859-43-4P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders)

100-39-0, Benzyl bromide 96-09-3, Styrene oxide 96-41-3, Cyclopentanol 100-51-6, Benzyl alcohol, reactions 100-46-9, Benzylamine, reactions 108-95-2, Phenol, reactions 107-08-4, 1-Iodopropane 110-89-4, Piperidine, reactions 110-91-8, Morpholine, reactions 111-49-9 122-60-1, 1,2-Epoxy-3-phenoxypropane 123-75-1, Pyrrolidine, reactions 288-36-8, 1H-1,2,3-Triazole 288-32-4, Imidazole, reactions 288-88-0, 288-94-8, 1H-Tetrazole 1H-1,2,4-Triazole 542-69-8, 1-Iodobutane 1855-36-3, 3,4-Dimethylstyrene oxide 2210-79-9, Glycidyl 1126-76-7 2211-94-1, Glycidyl 4-methoxyphenyl ether 2-methylphenyl ether 2212-05-7, 4-Chlorophenyl glycidyl ether 2783-26-8, 2-Methylstyrene 2788-86-5, 4-Chlorostyrene oxide 2783-28-0 3101-60-8, 4-tert-Butylphenyl glycidyl ether 5255-75-4, 4-Nitrophenyl glycidyl 6388-74-5, 4-Nitrostyrene oxide 13107-39-6, 4-Methylstyrene ether 13692-15-4, Oxirane, (2,4-dichlorophenyl)-18511-62-1, 20697-04-5, 3-Chlorostyrene oxide 4-Fluorostyrene oxide 20697-05-6, 3-Nitrostyrene oxide 20780-53-4, Oxirane, phenyl-, (2R)-20780-54-5, (S)-Styrene oxide 20861-99-8 21019-51-2, Oxirane, (4-chlorophenyl)-, 37586-22-4, 4-Benzoylpiperidine 52695-39-3 52909-94-1,

53220-41-0, 4-(4-Chlorobenzoyl)piperidine 3,4-Dichlorostyrene oxide 55967-94-7, 2-Oxiranylpyridine 56346-57-7, 4-(4-Fluorobenzoyl)piperidine 62717-50-4, 2-Chlorostyrene oxide 66256-03-9 71031-02-2 74130-04-4 76362-12-4, 4-(4-Methoxybenzoyl)piperidine 78038-42-3, (S)-4-Nitrostyrene oxide 78038-43-4, (R)-4-Nitrostyrene oxide 97466-49-4, (S)-4-Chlorostyrene oxide 93114-06-8 94829-51-3 111991-14-1, 4-Trifluoromethylstyrene oxide 111991-17-4 146145-08-6 169272-14-4 169272-15-5 478538-76**-**0 586417-77-8 666859-62-7 RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders) THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 9 (1) A Waverly co; Stedman's medical dictionary 1995, P362 (2) Anon; EP 409236 1991 HCAPLUS (3) Anon; Bundgaard Design of prodrugs 1986, P7 (4) Davis; US 4415581 A 1983 HCAPLUS (5) Gaudilliere; US 4711899 A 1987 HCAPLUS (6) Helsley; US 4812456 A 1989 HCAPLUS (7) Rae; US 5935974 A 1999 HCAPLUS (8) Rae; US 6365604 B1 2002 HCAPLUS (9) Wettlaufer; US 5114936 A 1992 HCAPLUS 666858-06-6P 666858-07-7P 666858-08-8P 666859-31-0P 666859-33-2P 666859-35-4P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (drug candidate; preparation of 4-benzoylpiperidine derivs. for treatment of psychosis and cognition disorders) 666858-06-6 HCAPLUS

HC1

666858-07-7 HCAPLUS RNMethanone, [1-[(2S)-2-[(aminocarbonyl)oxy]-2-phenylethyl]-4-piperidinyl](4-CN fluorophenyl) -, monohydrochloride (9CI) (CA INDEX NAME)

Methanone, [1-[2-[(aminocarbonyl)oxy]-2-phenylethyl]-4-piperidinyl](4-

fluorophenyl)-, monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE

IT

RN

CN

# ● HCl

RN 666858-08-8 HCAPLUS

CN Methanone, [1-[(2R)-2-[(aminocarbonyl)oxy]-2-phenylethyl]-4-piperidinyl](4-fluorophenyl)-, monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# HCl

RN 666859-31-0 HCAPLUS

CN Methanone, [1-[2-[(aminocarbonyl)oxy]-2-phenylethyl]-4-piperidinyl](4-fluorophenyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{Ph} & \text{O} \\ & | & | \\ \text{CH}_2-\text{CH}-\text{O}-\text{C}-\text{NH}_2 \\ \end{array}$$

RN 666859-33-2 HCAPLUS

CN Methanone, [1-[(2S)-2-[(aminocarbonyl)oxy]-2-phenylethyl]-4-piperidinyl](4-fluorophenyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 666859-35-4 HCAPLUS

CN Methanone, [1-[(2R)-2-[(aminocarbonyl)oxy]-2-phenylethyl]-4-piperidinyl](4-

# fluorophenyl) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

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L157 ANSWER 2 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
    1996:410460 HCAPLUS
AN
DN
    125:87211
    Entered STN: 16 Jul 1996
ED
    Preparation of O-(carbamoyl)phenylalaninol antidepressants
TI
    Choi, Yong Moon; Byun, Jai Kook
IN
    Yukong Limited, S. Korea
PA
    PCT Int. Appl., 26 pp.
SO
    CODEN: PIXXD2
DT
    Patent
LA
    English
    ICM C07C271-20
IC
    ICS C07C269-00
    34-2 (Amino Acids, Peptides, and Proteins)
CC
    Section cross-reference(s): 1, 25
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                                                               DATE
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                                                               19950906 <--
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                                          WO 1995-KR114
PΤ
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        W: AU, CA, CN, JP, RU, US
        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                                               19950906 <--
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    AU 700544
                        B2
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                                                               19950906 <--
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                              19991103
    EP 728129
                        _{\rm B1}
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
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                              19961106
    CN 1135209
                        Α
    CN 1069635
                        В
                              20010815
                                          JP 1995-509386
                                                               19950906 <--
                        T2
                              19970331
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                                                               19950906 <--
                                         RU 1996-113126
                              19980210
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    RU 2104266
                                         ES 1995-931444
                                                                19950906 <--
                              20000301
    ES 2140703
                        Т3
                                         US 1996-619657
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PRAI KR 1994-22798
                        Α
                              19940909 <--
                              19950906 <--
    WO 1995-KR114
                        W
CLASS
                CLASS PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
                _____
 ______
                ICM
                      C07C271-20
 WO 9607637
                      C07C269-00
                ICS
GΙ
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The title free-base compds. and their hydrochloride salts, I-III, useful in treating CNS diseases, particularly depression, are prepared by treating racemic, D-, or L-phenylalaninol with benzyl chloroformate in a basic aqueous solution to give the corresponding N-(benzyloxycarbonyl)phenylalaninol, reacting the intermediate with phosgene and then with an excess of a concentrated NH4OH aqueous solution to produce the corresponding O-carbamoyl-N-(benzyloxycarbonyl)phenylalaninol which is deprotected via hydrogenolysis, and the free base subjected to HCl salification. The free base of III was so prepared and demonstrated a 62% inhibition in the mouse forced-swimming depression model at 30 mg/kg (p.o.).

ST carbamoylphenylalaninol prepn antidepressant

IT Antidepressants

IT

(O-(carbamoyl)phenylalaninols)

178429-61-3P 178429-62-4P 178429-63-5P

178429-64-6P 178429-65-7P 178429-66-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation);

USES (Uses)

(preparation of O-(carbamoyl)phenylalinol antidepressants)

TT 75-44-5, Phosgene 501-53-1, Benzyl chloroformate 1336-21-6, Ammonium hydroxide 3182-95-4, L-Phenylalaninol 5267-64-1,

D-Phenylalaninol 7647-01-0, Hydrochloric acid, reactions

16088-07-6, Phenylalaninol

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of O-(carbamoyl)phenylalinol antidepressants)

IT 178429-61-3P 178429-62-4P 178429-63-5P

178429-64-6P 178429-65-7P 178429-66-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of O-(carbamoyl)phenylalinol antidepressants)

RN 178429-61-3 HCAPLUS

CN Benzenepropanol, β-amino-, carbamate (ester) (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{NH}_2 & \text{O} \\ & | & | \\ \text{Ph-CH}_2 - \text{CH-CH}_2 - \text{O-C-NH}_2 \end{array}$$

RN 178429-62-4 HCAPLUS

CN Benzenepropanol, β-amino-, carbamate (ester), (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$H_2N$$
  $O$   $R$   $Ph$ 

RN 178429-63-5 HCAPLUS

CN Benzenepropanol,  $\beta$ -amino-, carbamate (ester), (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 178429-64-6 HCAPLUS

CN Benzenepropanol,  $\beta$ -amino-, carbamate (ester), monohydrochloride (9CI) (CA INDEX NAME)

# ● HCl

RN 178429-65-7 HCAPLUS

CN Benzenepropanol,  $\beta$ -amino-, carbamate (ester), monohydrochloride, (R)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

#### HC1

RN 178429-66-8 HCAPLUS

CN Benzenepropanol,  $\beta$ -amino-, carbamate (ester), monohydrochloride, (S)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

# HCl

3182-95-4, L-Phenylalaninol 5267-64-1, D-Phenylalaninol IT 7647-01-0, Hydrochloric acid, reactions 16088-07-6, Phenylalaninol RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of O-(carbamoyl)phenylalinol antidepressants) 3182-95-4 HCAPLUS RN Benzenepropanol,  $\beta$ -amino-,  $(\beta S)$ - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

$$\Pr \bigcup_{NH_2}^{S} \mathsf{OH}$$

CN

5267-64-1 HCAPLUS RNBenzenepropanol, β-amino-, (βR)- (9CI) (CA INDEX NAME) CN

Absolute stereochemistry. Rotation (+).

7647-01-0 HCAPLUS RN Hydrochloric acid (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME) CN

HC1

16088-07-6 HCAPLUS RN Benzenepropanol,  $\beta$ -amino- (9CI) (CA INDEX NAME) CN

$$\begin{array}{c|c} & \mathrm{NH_2} \\ & | \\ \mathrm{HO-CH_2-CH-CH_2-Ph} \end{array}$$

L157 ANSWER 3 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN

1989:57520 HCAPLUS AN

DN 110:57520

Entered STN: 17 Feb 1989 ED

Preparation of N-containing heterocycles for treatment of cerebral ΤI disorders

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Sugimoto, Hachiro; Nakamura, Takaharu; Karibe, Norio; Saito, Isao;
IN
    Higurashi, Kunizo; Yonaga, Masahiro; Kaneko, Takeru; Nakazawa, Takahiro;
    Ueno, Masataka; Yamatsu, Kiyomi
PΑ
    Eisai Co., Ltd., Japan
    PCT Int. Appl., 53 pp.
SO
    CODEN: PIXXD2
DT
     Patent
LA
    Japanese
IC
     ICM C07D211-14
         C07D211-18; C07D211-22; C07D211-32; C07D211-70; C07D295-18;
          C07D401-06; C07D405-04; C07D409-04; A61K031-445; A61K031-47;
          A61K031-505
     27-16 (Heterocyclic Compounds (One Hetero Atom))
CC
     Section cross-reference(s): 1
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    WO 8802365
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         RW: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE
                                                                   19860930
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                         · A1
    AU 8664054
                                19900719
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    EP 288563
                          A1
                                19881102
                                            EP 1986-905925
                                19940511
     EP 288563
                          B1
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    NO 175055
PRAI EP 1986-905925
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     WO 1986-JP502
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CLASS
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                        PATENT FAMILY CLASSIFICATION CODES
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 WO 8802365
                 ICM
                        C07D211-14
                        C07D211-18; C07D211-22; C07D211-32; C07D211-70;
                 ICS
                        C07D295-18; C07D401-06; C07D405-04; C07D409-04;
                        A61K031-445; A61K031-47; A61K031-505
     CASREACT 110:57520; MARPAT 110:57520
OS
GΙ
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III

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Title compds. I [A = (substituted) Ph, pyridyl, thienyl, (substituted)
AB
     naphthyl, tetralyl, quinolyl, benzofuranyl, quinazolyl, benzothienyl,
     1-oxo-1,2,3,4-tetrahydronaphthalen-2-yl, 1,2,3,4-tetrahydronaphthoquinon-2-
     yl; X = CH2CO, CHOH, CHMe, CHCH2NEt2; Y = C, N; Z = CH2, CO, CHOR1 (R1 =
     H, alkyl, acyl, aralkyl, heteroaralkyl, CHR2 (R2 = halo), CH, p-R2C6H4C,
     CHR3 (R3 = N-succinimidyl); Z is bonded at the 3 or 4 position; B = halo,
     alkyl, alkoxy, (mono- or disubstituted) Ph, naphthyl; m = 1-3; n = 0-4;
     dashed line = double bond], useful for treatment and prevention of mental
     disorders induced by apoplexy, cerebrosclerosis, and cerebroinfarct, are
     prepared from heterocycles II. A mixture of 2-bromo-2'-acetonaphthone,
     4-(p-fluorobenzoyl)piperidine, HCl, KI and NaHCO3 in EtOH was refluxed to
     give III and III was converted to its HCl salt, which at 3 mg/kg p.o.
     showed 143% increase life span in ischemia-induced rats.
ST
     heterocycle nitrogen contg cerebral disorder; piperidine prepn treatment
     mental disorder
IT
    Mental disorder
        (treatment and prevention of, by nitrogen-containing heterocycles)
IT
     Brain, disease or disorder
        (cerebrovascular, treatment and prevention of, by nitrogen-containing
        heterocycles)
IT
     Brain, disease or disorder
        (ischemia, treatment and prevention of, by nitrogen-containing
        heterocycles)
IT
     95374-61-1P
                   107025-80-9P
                                  107025-81-0P
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                                                                 118411-69-1P
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     118412-23-0P
                    118412-29-6P
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                    118412-34-3P
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     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, for treatment and prevention of cerebral disorders)
                           2633-50-3
                                       5696-78-6
                                                    13686-51-6
IT
     613-54-7
                700-46-9
                                                                 20849-71-2,
     1-Chloro-2-(2-naphthyl)ethane
                                     31252-42-3, 4-Benzylpiperidine
                  56346-57-7, 4-(p-Fluorobenzoyl)piperidine
                                                               58113-36-3
     54924-33-3
                                                               118412-66-1
     92822-02-1, 4-(p-Fluorobenzyl)piperidine
                                                118412-65-0
     118412-67-2
                   118412-68-3
                                 118412-69-4
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, in preparation of drug for cerebral disorders)
IT
     118411-89-5P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, for treatment and prevention of cerebral disorders)
RN
     118411-89-5 HCAPLUS
    Methanone, (4-fluorophenyl) [1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-,
CN
    hydrochloride (9CI) (CA INDEX NAME)
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$$\begin{array}{c|c} & \text{Ph} & \\ & | \\ \text{C} & \\ \end{array}$$

#### HCl

GI

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L157 ANSWER 4 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
     1987:598088 HCAPLUS
AN
DN
     107:198088
     Entered STN: 27 Nov 1987
ED
     Preparation of 1-phenyl-2-(4-benzoylpiperidino)alkanols as cerebrovascular
TI
IN
     Gaudilliere, Bernard; Rousseau, Jean
     Synthelabo S. A., Fr.
PΑ
     Eur. Pat. Appl., 35 pp.
     CODEN: EPXXDW
DT
     Patent
LΑ
     French
IC
     ICM C07D211-32
     ICS A61K031-445
     27-16 (Heterocyclic Compounds (One Hetero Atom))
CC
     Section cross-reference(s): 63
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                        KIND
                               DATE
                                          APPLICATION NO.
                                                                 DATE
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                                        EP 1986-401000
    EP 202164
                         A1
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                         A1 19861121
                                           FR 1985-7270
                                                                 19850514
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                                          AU 1986-57406
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                         B2
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                               19890601
    ZA 8603533
                        Α
                               19861230
                                          ZA 1986-3533
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                        A2
                                          HU 1986-1975
    HU 40790
                               19870227
                                                                 19860513
                        В
    HU 195640
                               19880628.
                        A
    US 4711899
                                           US 1986-862715
                               19871208
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    ES 554895
                        A1
                                           ES 1986-554895
                               19880401
                                                                 19860513
    IL 78773
                         A1
                               19900917
                                          IL 1986-78773
                                                                 19860513
                               19880216
    ES 557582
                         A1
                                           ES 1987-557582
                                                                 19870608
PRAI FR 1985-7270
                               19850514
    EP 1986-401000
                               19860512
CLASS
 PATENT NO.
                CLASS
                       PATENT FAMILY CLASSIFICATION CODES
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                ICM
                       C07D211-32
EP 202164
                ICS
                       A61K031-445
os
    CASREACT 107:198088
```

The title compds. (I; R = H, Me; R1 = H, C1-4 alkyl, C1-4 alkoxy, OH, PhCH2O, CF3, cyano, NO2, NH2, NHAc, MeS, MeSO2, H2NSO2; R2 = H, F, C1, Me, MeO) were prepared as cerebrovascular agents (no data). Styrene oxide was refluxed 3 h with 4-(4-fluorobenzoyl)piperidine in MeOH containing K2O3 to give I (R = R1 = H, R2 = 4-F).

ST benzoylpiperidineethanol prepn cerebrovascular agent; piperidineethanol benzoyl prepn cerebrovascular agent; vasodilator cerebral benzoylpiperidineethanol prepn

Ι

IT Ischemia

(treatment of, phenylpiperidineethanols for)

IT Brain, disease or disorder

(ischemia, treatment of, phenylpiperidineethanols for)

IT Neurotransmitter agonists

(serotoninergic, phenylpiperdineethanols)

IT Receptors

RL: RCT (Reactant); RACT (Reactant or reagent) ( $\alpha$ 1-adrenergic, of cerebral cortex, binding to, by

phenylpiperidineethanols)
IT 99-03-6, 1-(3-Aminophenyl)ethanone

RL: RCT (Reactant); RACT (Reactant or reagent)
 (acetylation of)

IT 7463-31-2P, 1-[3-(Acetylamino)phenyl]ethanone

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and bromination of)

TT 111000-50-1P 111000-52-3P 111000-53-4P 111000-56-7P 111000-57-8P 111000-58-9P 111000-59-0P 111000-61-4P 111000-62-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and hydrolysis of)

IT 111000-49-8P 111000-51-2P 111000-55-6P 111000-60-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reduction of)

IT 30095-56-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and N-alkylation by, of dioxolanylpiperidine derivative)

110999-79-6P 110999-77-4P 110999-78-5P TΤ 109577-45-9P 110999-84-3P 110999-82-1P 110999-83-2P 110999-80-9P 110999-81-0P 110999-87-6P 110999-88-7P 110999-89-8P 110999-86-5P 110999-85-4P 110999-92-3P 110999-93-4P 110999-94-5P 110999-91-2P 110999-90-1P 110999-97-8P 110999-98-9P 110999-99-0P 110999-95-6P 110999-96-7P 111000-02-3P 111000-03-4P 111000-04-5P 111000-01-2P 111000-00-1P 111000-06-7P 111000-07-8P 111000-08-9P 111000-09-0P 111000-05-6P 111000-11-4P 111000-12-5P 111000-13-6P 111000-14-7P 111000-10-3P 111000-16-9P 111000-17-0P 111000-18-1P 111000-19-2P 111000-15-8P 111000-24-9P 111000-25-0P 111000-26-1P 111000-22-7P 111000-20-5P 111000-27-2P 111000-28-3P 111000-29-4P 111000-30-7P 111000-31-8P 111000-32-9P 111000-33-0P 111000-34-1P 111000-35-2P 111000-36-3P 111000-38-5P 111000-39-6P 111000-40-9P 111000-41-0P 111000-37-4P

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111000-47-6P
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     111000-42-1P
     111004-36-5P
                    111004-37-6P 111058-47-0P 111058-48-1P
     111058-49-2P 111058-50-5P 118411-89-5P
     145526-24-5P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, as cerebrovascular agent)
                              2788-86-5, 4-Chloro styrene oxide
IT
     96-09-3, Styrene oxide
                                                                   20697-03-4,
                              20780-53-4, (R)-(-)-Phenyloxirane
                                                                   20780-54-5,
     3-Methyl styrene oxide
     (S) - (+) -Phenyloxirane
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (N-alkylation by, of benzoylpiperidines)
     345-94-8 2632-13-5, 2-Bromo-1-(4-methoxyphenyl)ethanone
                                                                  2632-14-6
IT
     111000-54-5
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (N-alkylation by, of dioxolanylpiperidine derivative)
     25519-78-2, 4-(4-Fluorobenzoyl)piperidine hydrochloride
IT
                                                                56346-57-7,
     4-(4-Fluorobenzoyl)piperidine
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (N-alkylation of, by styrene oxides)
     53220-47-6
                111000-48-7
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (N-alkylation of, by \alpha-haloketones)
IT
     109577-45-9P 111058-47-0P 111058-48-1P
     111058-49-2P 111058-50-5P 118411-89-5P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, as cerebrovascular agent)
     109577-45-9 HCAPLUS
RN
CN
     Methanone, (4-fluorophenyl)[1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-
           (CA INDEX NAME)
```

RN 111058-47-0 HCAPLUS

CN Methanone, (4-fluorophenyl)[1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-, hydrochloride, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

## ● HCl

RN 111058-48-1 HCAPLUS

CN Methanone, (4-fluorophenyl) [1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-, hydrochloride, (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# HC1

RN 111058-49-2 HCAPLUS

CN Methanone, (4-fluorophenyl) [1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 111058-50-5 HCAPLUS

CN Methanone, (4-fluorophenyl) [1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-, (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 118411-89-5 HCAPLUS

CN Methanone, (4-fluorophenyl) [1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-, hydrochloride (9CI) (CA INDEX NAME)

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L157 ANSWER 5 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
    1987:477645 HCAPLUS
AN
    107:77645
DN
    Entered STN: 05 Sep 1987
ED
    Dihydropyridinedicarboxylates, procedure for their preparation, and their
TΙ
    use as cardiovascular agents
    Kuehnis, Hans
IN
    Ciba-Geigy A.-G. , Switz.
PΑ
    Eur. Pat. Appl., 32 pp.
SO
    CODEN: EPXXDW
DT
    Patent
T.A
    German
    ICM C07D211-90
IC
     ICS C07D409-12; A61K031-445
     27-16 (Heterocyclic Compounds (One Hetero Atom))
     Section cross-reference(s): 1
FAN.CNT 1
                                        APPLICATION NO.
                       KIND DATE
                                                              DATE
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                              _____
                                                                19861031
              A2 19880107
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        R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE
    DK 8605278 A 19870507 DK 1986-5278
AU 8664837 A1 19870514 AU 1986-64837
ZA 8608428 A 19870624 ZA 1986-8428
JP 62114965 A2 19870526 JP 1986-262896
                                                                19861105
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PRAI CH 1985-4759
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 EP 222702 ICM ICS
                      C07D211-90
                      C07D409-12; A61K031-445
GΙ
```

# \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Title compds. I (R = carbo- or heterocyclic aryl; R1 = alkyl; one of R2 AB and R3 = alkyl, the other = alkyl, cyano, NH2; X = 0, NH; Z = alkylenesubstituted with carbocyclic aryl, with X separated by  $\geq 2$  C atoms from ring N; Y = alkylene, CHOH, CO, bond; Ar1 = monocyclic aryl, heteroaryl) and their salts, useful as cardiovascular agents with Ca antagonistic and  $\alpha$ -receptor blocking activityand as coronary dilators and antihypertensives for treating cardiovascular disorders such as circulatory disorders, high blood pressure, arrhythmia, and heart insufficiency, were prepared by 5 methods, e.g. by ring closure of diene II (one of X' and Y' = NH2, the other OH or NH2) or a tautomer thereof. A mixture of 4-(4-fluorobenzoyl)piperidine, styrene oxide, and THF was refluxed 15 h to give 2-[4-(4-fluorobenzoyl)-1-piperidinyl]-1phenylethanol, which was esterified with the reaction product of (COC1)2 and the mono-Me ester of 1,4-dihydro-2,6-dimethyl-4-(3-nitrophenyl)-3,5pyridinedicarboxylic acid to give a diastereomeric mixture of diester III. The blood pressure of rats treated orally with 2.0 mg III/kg was lowered .apprx.70 mm after 2 h. The IC50 for III in in vitro testing on rat tissue was .apprx.5  $\mu$ mol/L, .apprx.4  $\mu$ mol/L, and .apprx.80  $\mu$ mol/L for K, noradrenaline, and serotonin induced vasoconstriction. Capsules containing 10 mg III were prepared from III 2500, talc 200, and colloidal silicic acid 50 mg.

ST coronary dilator pyridinedicarboxylate prepn; antihypertensive pyridinedicarboxylate prepn; circulation pyridinedicarboxylate prepn; calcium antagonist pyridinedicarboxylate prepn; serotonin antagonist

```
pyridinedicarboxylate prepn; noradrenaline antagonist
     pyridinedicarboxylate prepn; potassium antagonist pyridinedicarboxylate
     prepn; cardiovascular pyridinedicarboxylate prepn
    Antihypertensives
IT
        (pyridinedicarboxylate esters)
     Vasodilators
IT
        (coronary, pyridinedicarboxylate esters)
     50-67-9, Serotonine, biological studies
                                               51-41-2, Noradrenaline
TT
     7440-09-7, Potassium, biological studies 7440-70-2, Calcium, biological
     studies
     RL: BIOL (Biological study)
        (antagonists, pyridinedicarboxylate esters as)
                                109577-49-3
     74936-72-4 109577-47-1
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (esterification of)
IT
     109577-45-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and esterification of)
                                                                  109577-52-8P
                    109577-48-2P
                                                   109577-51-7P
                                   109577-50-6P
IT
     109577-46-0P
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                    109577-69-7P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (preparation of, as cardiovascular agent)
     96-09-3, Styrene oxide
TΤ
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with (fluorobenzoyl)piperidine)
     56346-57-7, 4-(4-Fluorobenzoyl)piperidine
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with styrene oxide)
TT
     109577-45-9P
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     (Reactant or reagent)
         (preparation and esterification of)
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RN
     Methanone, (4-fluorophenyl) [1-(2-hydroxy-2-phenylethyl)-4-piperidinyl]-
CN
     (9CI)
            (CA INDEX NAME)
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### => d l158 all fhitstr tot

L158 ANSWER 1 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:5766 HCAPLUS

DN 138:55858

ED Entered STN: 05 Jan 2003

TI Preparation of 2-heterocyclyl-1,2-ethanediol carbamates as nervous system agents.

IN Choi, Yong-Moon; Lee, Ki-Ho

PA SK Corporation, S. Korea

```
SO
     PCT Int. Appl., 33 pp.
    CODEN: PIXXD2
DT
     Patent
     English
LA
IC
     ICM A61K031-27
     27-8 (Heterocyclic Compounds (One Hetero Atom))
     Section cross-reference(s): 1, 28
FAN.CNT 1
                                           APPLICATION NO.
     PATENT NO.
                        KIND
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                               20030103
PΙ
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             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
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                                         EP 2002-741462
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            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                                                  20020621 <--
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                               20030424
                                          US 2002-177041
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PRAI US 2001-300730P
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     WO 2002-KR1147
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CLASS
                CLASS PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
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 WO 2003000247 ICM
                        A61K031-27
     MARPAT 138:55858
OS
     ACHB1CH2B2 [A = heterocyclyl optionally substituted by \geq 1 alkyl,
AB
     aryl, halo, trihalomethyl, trihalomethoxy, trialkylsilyl, SOR, SO2R,
     SO2NRR', SO3R, SR, NO2, NRR', OR, CN, COR10COR, NHCOR, CO2R, CONRR'; R, R'
     = H, alkyl, aryl; B1, B2 = OH, O2CNR1R2, provided that B1 and B2 are not
     simultaneously OH; R1, R2 = H, OH, alkyl, alkoxy, alkylaryl, arylalkyl,
     aryl, aryloxy], were prepared Thus, 1,1'-carbonyldiimidazole was added to a
     solution of 1-(2-thienyl)-1,2-ethanediol in CH2Cl2 at 5°; the reaction
     mixture was allowed to come to room temperature with stirring over 1 h aqueous
NH3 was
     added at 5° followed by stirring at room temperature for 1 h to give
     [2-(2-thieny1)-2-carbamoyloxyethyl]oxocarboxamide. Title compds.
     inhibited PTZ-induced convulsions in mice with ED50 = 31.3-50 mg/kg i.p.
     The compds. are effective in the treatment of disorders of the central
     nervous system, especially as anticonvulsive or antiepileptic agents.
     heterocyclylethanediol carbamate prepn anticonvulsant antiepileptic muscle
ST
     relaxant analgesic; thienylcarbamoyloxyethyloxocarboxamide prepn nervous
     system agent
IT
     Analgesics
     Anticonvulsants
     Nervous system agents
        (preparation of 2-heterocyclyl-1,2-ethanediol carbamates as nervous system
        agents)
IT
     Muscle, disease
        (spasm, treatment; preparation of 2-heterocyclyl-1,2-ethanediol carbamates
        as nervous system agents)
IT
     Muscle relaxants
        (spasmolytics; preparation of 2-heterocyclyl-1,2-ethanediol carbamates as
        nervous system agents)
IT
     Brain, disease
        (stroke, treatment; preparation of 2-heterocyclyl-1,2-ethanediol carbamates
        as nervous system agents)
```

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IT
     Pain
        (treatment; preparation of 2-heterocyclyl-1,2-ethanediol carbamates as
        nervous system agents)
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IT
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     RL: PAC (Pharmacological activity); SPN (Synthetic preparation);
     THU (Therapeutic use); BIOL (Biological study); PREP (Preparation)
     ; USES (Uses)
        (preparation of 2-heterocyclyl-1,2-ethanediol carbamates as nervous system
        agents)
     74-89-5, Methylamine, reactions
                                       530-62-1, 1,1'-Carbonyldiimidazole
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     3944-00-1, 1-(2-Pyridyl)-1,2-ethanediol
                                               19377-75-4,
     1-(2-Furanyl)-1,2-ethanediol 52098-28-9, 1-(2-Indolyl)-1,2-
                 67162-00-9, 1-(4-Methyl-5-thiazolyl)-1,2-ethanediol
     ethanediol
                                                 479639-87-7,
     143314-50-5, 1-(2-Thienyl)-1,2-ethanediol
     (+)-(1R)-1-(2-Thienyl)-1,2-ethanediol
                                             479639-88-8, (-)-(1S)-1-(2-
     Thienyl)-1,2-ethanediol 479639-89-9, 1-(5-Chloro-2-thienyl)-1,2-
                 479639-90-2, (+)-(1R)-1-(5-Chloro-2-thienyl)-1,2-ethanediol
     ethanediol
     479639-91-3, (-)-(1S)-1-(5-Chloro-2-thienyl)-1,2-ethanediol
                                                                  479639-92-4,
                                            479639-93-5, 1-(3,4,5-Trichloro-2-
     1-(5-Phenyl-2-thienyl)-1,2-ethanediol
     thienyl) -1,2-ethanediol 479639-94-6, 1-(5-Methyl-2-thienyl) -1,2-
                  479639-95-7, 1-(2,5-Dichloro-3-thienyl)-1,2-ethanediol
     ethanediol
                                                          479639-97-9
     479639-96-8, 1-(3-Chloro-2-thienyl)-1,2-ethanediol
     479639-98-0, 1-(5-Trifluoromethyl-2-thienyl)-1,2-ethanediol
                                                                    479639-99-1,
     1-(5-tert-Butyl-2-thienyl)-1,2-ethanediol
                                                 479640-00-1,
                                            479640-01-2, 1-(5-Trimethylsilyl-2-
     1-(5-Cyano-2-thienyl)-1,2-ethanediol
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     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of 2-heterocyclyl-1,2-ethanediol carbamates as nervous system
        agents)
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     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of 2-heterocyclyl-1,2-ethanediol carbamates as nervous system
        agents)
              THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
RE
(1) Anon; EUR POLYM J 1993, V29(9), P1217
(2) Anon; J MED CHEM 1967, V10(3), P491
(3) Anon; NOUV J CHIM 1978, V2(2), P119
(4) Forschungsinstitut Borstel Institut Fur Experimentelle; US 5798343 1998
    HCAPLUS
(5) Milliken Research Corporation; EP 918057 A 1999 HCAPLUS
     479639-79-7P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation);
     THU (Therapeutic use); BIOL (Biological study); PREP (Preparation)
     ; USES (Uses)
        (preparation of 2-heterocyclyl-1,2-ethanediol carbamates as nervous system
        agents)
     479639-79-7 HCAPLUS
RN
     1,2-Ethanediol, 1-(2-pyridinyl)-, bis(carbamate) (ester) (9CI)
                                                                      (CA INDEX
CN
     NAME)
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L158 ANSWER 2 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
    1998:352653 HCAPLUS
AN
DN
    129:28207
    Entered STN: 11 Jun 1998
ED
    Preparation of O-carbamoylphenylalaninol compounds as central nervous
TI
    system agents
    Choi, Yong Moon; Han, Dong Il; Kim, Yong Kil; Shin, Hun Woo;
IN
    Park, Jeong-han
    Yukong Ltd., S. Korea
PA
    U.S., 15 pp., Cont.-in-part of U.S. 5,705,640.
so
    CODEN: USXXAM
DT
    Patent
LA
    English
    ICM C07C261-00
IC
NCL
    560115000
    34-2 (Amino Acids, Peptides, and Proteins)
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                                         APPLICATION NO.
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PΙ
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                       Α
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                       C1 20010320
    US 5705640
PRAI KR 1995-2543
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                            19950211 <--
    US 1996-596496
                       A2 19960205 <--
CLASS
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 US 5756817
               ICM
                      C07C261-00
               NCL
                      560115000
    MARPAT 129:28207
OS
     Title compds. ArCH2CH(NH2)CH2OCONR1R2 (Ar = (substituted) phenyl; R1, R2 =
AB
    H, lower alkyl, aryl, arylalkyl, cyclopropyl, cyclohexyl; R1R2 =
     pyrrolidino, piperidino, morpholino, 4-methyl- or 4-phenylpiperazino,
     etc.) and their pharmaceutically useful salts were prepared The D-isomers
     of the title compds. were also prepared Title compds. are useful as central
     nervous system agents (no data).
     carbamoylphenylalaninol deriv racemic enantiomeric prepn; nervous system
ST
     agent carbamoylphenylalaninol deriv prepn
IT
     Nervous system agents
        (preparation of O-carbamoylphenylalaninol compds. as central nervous system
       agents)
     463-77-4DP, Carbamic acid, ester with phenylalaninol derivs., preparation
IT
     181797-92-2P 181797-93-3P 181797-94-4P
     181797-95-5P 181797-96-6P
                              181797-97-7P
                                             181797-98-8P
     181797-99-9P
                 181798-00-5P 183668-91-9P 183668-93-1P
     183668-95-3P 183668-97-5P 183668-99-7P
     183669-01-4P 183669-02-5P 183669-03-6P
     183669-04-7P 183669-05-8P 183669-06-9P
     183669-07-0P 183669-08-1P 183669-09-2P
     183669-10-5P 206063-38-9P
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    208119-28-2P 208119-29-3P
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation);
     USES (Uses)
        (preparation of O-carbamoylphenylalaninol compds. as central nervous system
        agents)
     58917-85-4, N-Benzyloxycarbonyl-D-phenylalaninol
IT
     183669-11-6 183669-12-7 183669-14-9
     183669-15-0 206063-99-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of O-carbamoylphenylalaninol compds. as central nervous system
        agents)
     181797-75-1P 181797-77-3P 181797-78-4P
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                                                 181797-87-5P
     181797-79-5P 181797-82-0P
                    181797-91-1P 183668-80-6P 183668-83-9P
     181797-89-7P
     183668-85-1P 183668-87-3P 183668-89-5P
     183669-13-8P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (preparation of O-carbamoylphenylalaninol compds. as central nervous system
        agents)
     181797-92-2P
IT
     RL: BAC (Biological activity or effector, except adverse); SPN
     (Synthetic preparation); PREP (Preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation);
     USES (Uses)
        (preparation of O-carbamoylphenylalaninol compds. as central nervous system
        agents)
RN
     181797-92-2
                 HCAPLUS
     Benzenepropanol, β-amino-, methylcarbamate (ester),
CN
     monohydrochloride, (βR) - (9CI) (CA INDEX NAME)
Absolute stereochemistry.
MeNH
               NH<sub>2</sub>
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HC1

L158 ANSWER 3 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
AN 1998:239197 HCAPLUS
DN 128:295052
ED Entered STN: 27 Apr 1998
TI Preparation of O-carbamoyl-phenylalaninol compounds and their

```
pharmaceutically useful salts
    Choi, Yong Moon; Han, Dong Il; Kim, Yong Kil; Shin, Hun Woo;
IN
    Park, Jeong Han
    Yukong Limited, S. Korea
PΑ
SO
    PCT Int. Appl., 54 pp.
    CODEN: PIXXD2
DT
    Patent
    English
LΑ
IC
     ICM C07C271-12
     34-2 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 1
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                                           APPLICATION NO.
                                                                  DATE
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                               19980416
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 WO 9815526
                       C07C271/20; C07C271/24; C07C271/28; C07D295/20B5
                ECLA
 CA 2240060
   MARPAT 128:295052
OS
    Racemic or enantiomerically enriched O-carbamoyl-phenylalaninol compds.
AB
     PhCH2CH(NH2)CH2O2CNR1R2 (Ph may be substituted; R1, R2 = H, alkyl, aryl,
     arylalkyl, cyclic Pr, cycloaliph. or R1R2N is a cyclic group which may
     contain an addnl. nitrogen atom which may be substituted or an oxygen
     atom) or their pharmaceutically acceptable salts were prepared Thus,
     O-carbamoyl-o-fluorophenylalaninol hydrochloride was prepared from
     N-(tert-butoxycarbonyl)-o-fluorophenylalaninol by treatment with
     1,1'-carbonyldiimidazole in THF and then ammonia and deprotection by 6N
     carbamoyl phenylalaninol prepn pharmaceutical
ST
IT
        (preparation of O-carbamoylphenylalaninol compds. and their pharmaceutically
        useful salts)
     92-54-6, n-Phenylpiperazine 108-91-8, Cyclohexanamine, reactions
IT
     110-89-4, Piperidine, reactions 110-91-8, Morpholine, reactions
     111-86-4, Octylamine 123-75-1, Pyrrolidine, reactions 58917-85-4
     183669-11-6 183669-12-7 183669-13-8
     183669-14-9 183669-15-0 206063-99-2
                              206064-05-3
     206064-00-8 206064-03-1
     206064-07-5 206064-08-6 206064-09-7
     206064-10-0 206064-11-1 206064-13-3
     206064-15-5 206064-17-7
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of O-carbamoylphenylalaninol compds. and their pharmaceutically
        useful salts)
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     183668-85-1P 183668-87-3P 183668-89-5P
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RL: RCT (Reactant); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation);
     RACT (Reactant or reagent); USES (Uses)
        (preparation of O-carbamoylphenylalaninol compds. and their pharmaceutically
        useful salts)
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     206064-21-3P 206064-26-8P
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (preparation of O-carbamoylphenylalaninol compds. and their pharmaceutically
        useful salts)
              THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
RE
(1) Berger, F; US 2937119 A 1960 HCAPLUS
(2) Yukong Limited; WO 9624577 A1 1996 HCAPLUS
     58917-85-4
     RL: SPN (Synthetic preparation); SPN (Synthetic
     preparation); PREP (Preparation); PREP
     (Preparation)
        (preparation of O-carbamoylphenylalaninol compds. and their pharmaceutically
        useful salts)
     58917-85-4 HCAPLUS
RN
     Carbamic acid, [(1R)-1-(hydroxymethyl)-2-phenylethyl]-, phenylmethyl ester
CN
            (CA INDEX NAME)
     (9CI)
Absolute stereochemistry.
             OH
Ph
L158 ANSWER 4 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
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DN 126:7835
ED Entered STN: 06 Dec 1996
TI O-Carbamoyl-phenylalaninol having substituent at benzene ring, its pharmaceutically useful salts and method for preparing the same
IN Choi, Yong Moon; Han, Dong Il; Kim, Yong Kil; Shin, Hun Woo

ΑN

1996:716301 HCAPLUS

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PΑ
    Yukong Limited, S. Korea
     PCT Int. Appl., 30 pp.
SO
    CODEN: PIXXD2
DT
    Patent
    English
LΑ
IC
    ICM C07C271-10
     ICS C07C269-00
CC
     25-21 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
    Section cross-reference(s): 1, 34
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                                                          19960409 <--
    CA 2217771
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                              19961017
                                         CA 1996-2217771
                                                               19960410 <--
    EP 820438
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                                         EP 1996-909388
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                        В1
    EP 820438
                              20010620
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    CN 1071741
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CLASS
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WO 9632375
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                ICS
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US 6140532
               ECLA
                      C07C271/12; C07C323/32
    MARPAT 126:7835
OS
GI
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$$O$$
 $NH_2$ 
 $O$ 
 $NH_2$ 

AB Title compds. I and pharmaceutically acceptable salts are disclosed [wherein R = C1-8 alkyl, halo, C1-3 alkoxy or alkylthio, NO2, OH, or CF3; x = 1-3, multiple R's may vary when x = 2 or 3]. Uses of I include treatment and prophylaxis of CNS disorders such as depression, anxiety, epilepsy, stroke, dementia, and Parkinson's disease (no data). For instance, N-(tert-butoxycarbonyl)-o-fluorophenylalaninol in THF was treated with 1,1'-carbonyldiimidazole and then NH3 to give 75% of the O-carbamoyl derivative This was deprotected with HCl in aqueous THF, and the product was acidified with anhydrous HCl in THF and precipitated with Et2O, to give

73% title compound 2-FC6H4CH2CH(NH2)CH2OCONH2.HCl.

I

ST carbamoylphenylalaninol prepn CNS agent; phenylalaninol carbamoyl prepn antidepressant anxiolytic

```
TΤ
    Mental disorder
        (dementia, treatment; preparation of carbamoylphenylalaninols as CNS agents)
TT
     Anticonvulsants
     Antidepressants
     Anxiolytics
     Cognition enhancers
     Nervous system agents
        (preparation of carbamoylphenylalaninols as CNS agents)
IT
     Brain, disease
        (stroke, treatment; preparation of carbamoylphenylalaninols as CNS agents)
IT
     Parkinson's disease
        (treatment; preparation of carbamoylphenylalaninols as CNS agents)
     183668-80-6P, O-Carbamoyl-N-(tert-butoxycarbonyl)-o-
IT
     fluorophenylalaninol 183668-83-9P, O-Carbamoyl-N-(tert-
     butoxycarbonyl)-p-fluorophenylalaninol 183668-85-1P,
     O-Carbamoyl-N-(tert-butoxycarbonyl)-p-nitrophenylalaninol
     183668-87-3P, O-Carbamoyl-N-(tert-butoxycarbonyl)-p-[(tert-
     butoxycarbonyl)oxy]phenylalaninol 183668-89-5P,
     O-Carbamoyl-N-[(benzyloxy)carbonyl]-m-fluorophenylalaninol
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP
     (Preparation); RACT (Reactant or reagent)
        (intermediate; preparation of carbamoylphenylalaninols as CNS agents)
IT
     183668-91-9P, O-Carbamoyl-o-fluorophenylalaninol hydrochloric acid
     salt 183668-93-1P, O-Carbamoyl-p-fluorophenylalaninol
     hydrochloric acid salt 183668-95-3P, O-Carbamoyl-p-
     nitrophenylalaninol hydrochloric acid salt 183668-97-5P,
     O-Carbamoyl-p-hydroxyphenylalaninol hydrochloric acid salt
     183668-99-7P, O-Carbamoyl-m-fluorophenylalaninol hydrochloric acid
     salt 183669-01-4P, O-Carbamoyl-o-fluorophenylalaninol
     183669-02-5P, O-Carbamoyl-p-chlorophenylalaninol
     183669-03-6P, O-Carbamoyl-m-fluorophenylalaninol
     183669-04-7P, O-Carbamoyl-p-nitrophenylalaninol
     183669-05-8P, O-Carbamoyl-p-fluorophenylalaninol
     183669-06-9P, O-Carbamoyl-p-(methylthio)phenylalaninol
     183669-07-0P, O-Carbamoyl-p-hydroxyphenylalaninol
     183669-08-1P, O-Carbamoyl-p-methoxyphenylalaninol
     183669-09-2P, O-Carbamoyl-3,4-dihydroxyphenylalaninol
     183669-10-5P, O-Carbamoyl-3,4-dimethoxyphenylalaninol
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation);
     USES (Uses)
        (preparation of carbamoylphenylalaninols as CNS agents)
     530-62-1, 1,1'-Carbonyldiimidazole
                                         7664-41-7, Ammonia, reactions
IT
     183669-11-6, N-(tert-Butoxycarbonyl)-o-fluorophenylalaninol
     183669-12-7, N-(tert-Butoxycarbonyl)-p-fluorophenylalaninol
     183669-13-8, N-(tert-Butoxycarbonyl)-p-nitrophenylalaninol
     183669-14-9, N-(tert-Butoxycarbonyl)-p-[(tert-
     butoxycarbonyl)oxy]phenylalaninol 183669-15-0,
    N-[(Benzyloxy)carbonyl]-m-fluorophenylalaninol
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (starting material; preparation of carbamoylphenylalaninols as CNS agents)
IT
     183668-80-6P, O-Carbamoyl-N-(tert-butoxycarbonyl)-o-
     fluorophenylalaninol
     RL: RCT (Reactant); SPN (Synthetic preparation); SPN
     (Synthetic preparation); RACT (Reactant or reagent); PREP
     (Preparation)
        (intermediate; preparation of carbamoylphenylalaninols as CNS agents)
RN
     183668-80-6 HCAPLUS
CN
    Carbamic acid, [2-[(aminocarbonyl)oxy]-1-[(2-fluorophenyl)methyl]ethyl]-,
     1,1-dimethylethyl ester (9CI) (CA INDEX NAME)
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L158 ANSWER 5 OF 5 HCAPLUS COPYRIGHT 2004 ACS on STN
    1996:605524 HCAPLUS
ΑN
    125:248474
DN
    Entered STN: 11 Oct 1996
ED
TI
    Preparation of O-carbamoyl-D-phenylalaninol CNS agents
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IN
PA
    Yukong Limited, S. Korea
SO
    PCT Int. Appl., 31 pp.
    CODEN: PIXXD2
DT
    Patent
LA
    English
IC
    ICM C07C271-12
    ICS C07C269-04; C07C295-205
CC
    34-2 (Amino Acids, Peptides, and Proteins)
    Section cross-reference(s): 28
FAN.CNT 2
                       KIND
    PATENT NO.
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ΡI
    WO 9624577
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                              19960815
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        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
    CA 2212326
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                                      CA 1996-2212326
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                              19980107
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PRAI KR 1995-2543
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    WO 1996-KR18
CLASS
               CLASS PATENT FAMILY CLASSIFICATION CODES
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WO 9624577
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                      C07C271-12
               ICS
                      C07C269-04; C07C295-205
OS
    MARPAT 125:248474
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AB 800-carbamoyl-(D)-phenylalaninols [I; R1, R2 = H, C1-8 alkyl, (un)substituted cycloaliph. heterocyclyl; the number of C atoms in both R1 and R2 is 0-16], useful as CNS agents (no data) in the treatment of depression (no data), anxiety (no data), epilepsy (no data), etc. (no

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data), are prepared by the reaction of D-phenylalaninol with benzyl
chloroformate, followed by carbamoylation of the protected aminoalc. with
phosgene, followed by amidation of the carbonate chloride with amines
R1(R2)NH. Thus, N-benzyloxycarbonyl-D-phenylalaninol was carbamoylated
with phosgene and the intermediate amidated with H2NMe, producing I (R1 =
H, R2 = Me) in 78\% yield.
carbamoylphenylalaninol prepn CNS agent; anxiolytic prepn
carbamoylphenylalaninol; antidepressant prepn carbamoylphenylalaninol;
antiepileptic prepn carbamoylphenylalaninol
Analgesics
Anticonvulsants and Antiepileptics
Antidepressants
Anxiolytics
Nervous system agents
   (O-carbamoyl-D-phenylalaninols)
                                  50-81-7, Ascorbic acid, reactions
50-21-5, Lactic acid, reactions
                                   60-80-0, Antipyrine
56-84-8, Aspartic acid, reactions
                                65-85-0, Benzoic acid, reactions
64-19-7, Acetic acid, reactions
69-72-7, reactions 75-31-0, Isopropylamine, reactions
                                                          75-44-5,
Carbonic dichloride 75-75-2, Methanesulfonic acid
                                                    75-92-3,
Hydroxymethanesulfonic acid 77-92-9, Citric acid, reactions
Tartaric acid, reactions 92-54-6, N-Phenylpiperazine 98-11-3,
                                107-36-8
                                           108-91-8, Cyclohexylamine,
Benzenesulfonic acid, reactions
                                                 110-16-7, Maleic acid,
            110-15-6, Succinic acid, reactions
reactions
                                                110-89-4, Piperidine,
            110-17-8, Fumaric acid, reactions
reactions
                                            111-86-4, 1-Aminooctane
            110-91-8, Morpholine, reactions
reactions
121-44-8, Triethylamine, reactions
                                    121-69-7, Dimethylphenylamine,
            123-75-1, Pyrrolidine, reactions
                                               141-82-2, Propanedioic
reactions
acid, reactions 144-62-7, Oxalic acid, reactions
                                                     501-53-1, Benzyl
              526-95-4, Gluconic acid 594-45-6, Ethanesulfonic
chloroformate
      3424-21-3, Triisopropylamine 5267-64-1, D-Phenylalaninol
               6915-15-7, Malic acid
                                       7087-68-5, Diisopropylethylamine
6674-22-2, DBU
7647-01-0, Hydrochloric acid, reactions 7664-38-2,
Phosphoric acid, reactions 7664-93-9, Sulfuric acid, reactions
58917-85-4
RL: RCT (Reactant); RACT (Reactant or reagent)
   (preparation of O-carbamoyl-D-phenylalaninol CNS agents)
181797-75-1P 181797-77-3P 181797-78-4P
181797-79-5P 181797-82-0P
                            181797-84-2P
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181797-89-7P
181797-94-4P 181797-95-5P 181797-96-6P
                              181797-99-9P
                                             181798-00-5P
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181798-04-9P 181798-05-0P
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181798-07-2P
               181798-08-3P 181798-09-4P
               181798-12-9P 181798-13-0P
181798-11-8P
181798-14-1P
RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
   (preparation of O-carbamoyl-D-phenylalaninol CNS agents)
56-84-8, Aspartic acid, reactions
RL: SPN (Synthetic preparation); RACT (Reactant or reagent);
PREP (Preparation)
   (preparation of O-carbamoyl-D-phenylalaninol CNS agents)
56-84-8 HCAPLUS
L-Aspartic acid (9CI) (CA INDEX NAME)
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Absolute stereochemistry. Rotation (+).

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